

EUROPEAN PATENT SPECIFICATION

(45) Date of publication and mention of the grant of the patent: 06.05.1998 Bulletin 1998/19

(21) Application number: 94303969.3

(22) Date of filing: 02.06.1994

(12)

(51) Int CL6: C12N 9/24, C12N 9/90. C12P 19/12, C12P 19/14, C12P 19/24, A61K 31/70, A61K 7/00, A23L 1/236 // C12N9:24,(C12R1/06, 1:13, 1:265, 1:41)

(54) Trehalose-releasing enzyme, and its preparation and uses

Trehalose-freisetzendes Enzym, seine Herstellung und Verwendung Enzyme liberant du tréhalose, sa préparation et utilisation

(84) Designated Contracting States: AT BE CHIDE DK ES FRIGBIT LIMONL PT SE

(30) Priority: 03.06.1993 JP 156338/93 09.12.1993 JP 340343/93 28.03.1994 JP 79291/94

(43) Date of publication of application: 14.12.1994 Bulletin 1994/50

(73) Proprietor: KABUSHIKI KAISHA HAYASHIBARA SEIBUTSU KAGAKU KENKYUJO Okayama-shi Okayama (JP)

(72) Inventors:

 Maruta, Kazuhiko Okayama-shi, Okayama (JP) · Kubota, Michio Ibaraki-shi, Osaka (JP)

· Sugimoto, Toshiyuki

(11)

Okayama-shi, Okayama (JP)

· Miyake, Toshi Okayama-shi, Okayama (JP)

(74) Representative: Daniels, Jeffrey Nicholas et al Page White & Farrer 54 Doughty Street

London WC1N 2LS (GB) (56) References cited: FR-A- 2 671 099

EP-A- 0 483 755

EP-A- 0 555 540

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

Description

15

25

35

40

45

55

The present invention relates to a trehalose-releasing enzyme, and its preparation and uses, more particularly, to a novel trehalose-releasing enzyme which specifically hydrolyzes the linkage between a trehalose motely and the remaining glycosyl moiety in non-reducing saccharides having a trehalose structure as an end unit and having a glucose polymerization degree of 3 or higher, and to the preparation of the enzyme. The present invention further relates to microorganisms capable of forming the enzyme, trehalose obtained by using the enzyme, and compositions containing the trehalose.

Trehalose or α , α -trehalose has been known as a non-reducing seacharide consisting of glucose units. As is described in Advances in Carbohydrate Chemistry, Vol.18, pp.201-225 (1963), published by Academic Press, USA, and <math>Applied and Environmental Microbiology, Vol.56, pp.2.213-3.215 (1990), trehalose widely exists in microorganisms, mushrooms, insects, etc., though the content is relatively low. Trehalose is a non-reducing seacharide, so that it neither reacts with substances containing amino groups such as amino acids and proteins, induces the amino-carbonyl reaction, nor deteriorates amino acid-containing substances. Thus, trehalose would be used without a fear of causing an unsatisfactory browning and deterioration. Because of these, the establishment of the industrial-scale preparation of trehalose has been in oreal clampad.

Conventional preparations of trahalose are, for example, those which are disclosed in Japanese Patent Laid-Open No.154.68576 whereir microorganisms are utilized, and reported in Japanese Patent Laid-Open No.154.685763 wherein maltose is converted into trahalose by using maltose- and trahalose-phosphorylases in combination. The former, however, is not suitable for the industrial-scale preparation because the content of trahalose present in microorganisms used as a starting material is usually lower than 15 wW % (the wording "ww %" is abbreviated as "5" in the specification, unless otherwise specified), on a dry solid basis (d.s.b.), and the extraction and purification stops are complicated. The latter has the following demerklis Since trahalose is formed via glucose-l-phosphate, the concentration of maltose as a substrate could not be set to a satisfactority high-level; (ii) the enzymatic reaction systems of the phosphorylases are reversible reactions, and their yields of the objective trahalose are relatively low; and (iii) it is substantially difficult to retain their reaction systems stably and to continue their enzymatic reactions smoothly. Thus, those conventional preparations have not been actually used as an industrial-scale preparation.

As regards the preparation of trehalose, it is reported in the column titled "Oligoseccharides" in the chapter titled "Current Status of Starch Application Development and Related Problems" in "Food Chemicals", No 89, pp. 87-72 (August, 1992) that "In spile of a wide applicability of trehalose, the enzymatic preparation via a dired saccharide-transfer reaction or a hydrolytic reaction has been reported to be scientifically aimost impossible in this field." Thus, the prepration of trehalose by an enzymatic reaction sung starch as a material has ben deemed to be scientifically very difficult.

It is known that partial starch hydrolysates, prepared from a material starch such as liquefled starch, dextrins and maltooligosaccharides, exhibit a reducing power owing to their reducing and groups. The reducing power of these reducing partial starch hydrolysates is generally expressed by "Dextrose Equivalent (DE) value", based on a dry weight. It is known that among reducing partial starch hydrolysates those with a relatively-high DE value generally have a relatively-on noticular weight and viscosity, as well as a relatively-high level of sweetness and reactivity, and readily react with substances having amino groups such as amino acids and proteins to cause an unsatisfactory browning, smell and deletrioration of their quality.

Since the properties of reducing partial starch hydrolysates are varied dependently on their DE values, the relationship between reducing partial starch hydrolysates and their DE values is significant. It has been even believed impossible to break away the relationship in this field.

The present inventors, however, did change this common sense and succeeded to establish a preparation of trehalose as disclosed in Japanese Patent Application No.382,131/92 wherein trehalose is directly produced from non-reducing partial starch hydrolystates by allowing glucoamylase together with a non-reducing saccharide-forming enzyme capable of forming non-reducing saccharides, having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher, to act on reducing partial starch hydroystate having a degree of glucose polymerization of 3 or higher, prepared from a material starch. Although the preparation of trehalose yields trehalose from non-reducing partial starch hydrolystates in a yield of about 30% and can be feasible as an industrial-scale preparation, there still remains some fear of resulting in a high production cost in view of the trehalose yield. Therefore, the establishment of a novell preparation of trehalose, which forms trehalose from non-reducing partial starch hydrolysates in an increased yield is in great demand.

The object of the present invention is to provide a novel preparation of trehalose in a relatively-high yield from relatively-low cost and stably suppliable starch.

In order to attain the aforementioned object, the present inventors have extensively screened microorganisms capable of producing a novel enzyme which releases trehalose from non-reducing partial starch hydrolysates having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher. As a result, we found that a non-reducing saccharide-forming microorganism of the genus *Rhizobium*, i.e. *Rhizobium* sp. M-11, isolated from

a solid, as disclosed in Japanese Patent Application No.362.131/92, and a non-reducing saccharide-forming microcryanism of the genus Arthrobactor, ic Arthrobactor sp. 0.36, isolated from a soil, as disclosed in Japanese Patent Application No.265.416/93, produces a novel trehalose-releasing enzyme. We also tound that the novel trehalose-releasing enzyme taclitates a reaction to form trehalose in a satisfactorily-high pitod when used in combination with a non-reducing saccharide-forming enzyme, and that trehalose is readily prepared by allowing the novel trehalose-releasing enzyme and a non-reducing saccharide-forming enzyme to act on reducing partial starch hydrolysates, and recovering a reaction mixture containing a relatively-high purity trehalose. We extensively screened microorganisms which produce such a trehalose-releasing enzyme from among known microorganisms. As a result we found that microorganisms of the genera Brevibacterium and Microoccus produce a trehalose-releasing enzyme which forms trehalose from non-reducing saccharides having a trehalose-structure as an end unit and having a diegree of glucose polymerization of 3 or higher similarly as the trehalose-releasing enzymes derived from microorganisms of the genera Brizobium and Arthrobactor, and accomplished this invention. We also established compositions such as food products, cosmelics and pharmaceuticals containing the trehalose prepared by the aforesaid preparation, and accomplished this invention.

The present invention will now be described in further detail by way of example only. with reference to the accompanying drawings, in which:

FIG.1 shows elution patterns of the present trehalose-releasing enzyme and a non-reducing saccharide-forming enzyme eluted from a column packed with a gel of "DEAE-Toyopearl®".

FIG.2 shows the influence of temperature on the activity of the present trehalose-releasing enzyme derived from a microorganism of the species *Rhizobium* sp. M-11.

FIG.3 shows the influence of pH on the activity of the present trehalose-releasing enzyme derived from a microorganism of the species *Rhizobium* sp. M-11.

FIG.4 shows the influence of temperature on the stability of the present trehalose-releasing enzyme derived from a microorganism of the species *Rhizobium* sp. M-11.

FIG.5 shows the influence of pH on the stability of the present trehalose-releasing enzyme derived from a microorganism of the species *Rhizobium* sp. M-11.

FIG.6 shows the influence of temperature on the activity of the present trehalose-releasing enzyme derived from a microorganism of the species Arthrobacter sp. Q36.

FIG.7 shows the influence of pH on the activity of the present trehalose-releasing enzyme derived from a microorganism of the species Arthrobacter sp. Q36.

FIG.8 shows the influence of temperature on the stability of the present trehalose-releasing enzyme derived from a microorganism of the species *Arthrobacter* sp. Q36.

FIG.9 shows the influence of pH on the stability of the present trehalose-releasing enzyme derived from a microorganism of the species Arthrobacter so. Q36.

The identification test of a microorganism of the genus *Rhizobium*, i.e. "*Rhizobium* sp. M-11" according to the present invention gave the following results. The test was conducted in accordance with the method as described in "*Biseibutsu-no-Bunnui-to-Dotel*" (Classification and Identification of Microorganisms), edited by Takeji Hasegawa, published by Japan Scientific Societies Press, Tokyo, Japan (1985):

A. Morphology

15

20

25

30

35

45

50

55

Characteristics of cells when incubated at 27°C in nutrient agar

Usually existing in a rod form of 0.6-0.8x1.0-1.5µm;

Existing single but uncommonly existing in a serially coupled- or a linked-form;

Exhibiting no polymorphism;

Possessing motility, asporogenicity and flagellum;

Non-acid fast:

Gram stain : Negative;

Capsule : Negative:

Metachromatic granule: Positive; and

Accumulating poly-β-hydroxy butyrate.

B. Cultural property

(1) Characteristics of colony formed when incubated at 27°C in nutrient agar plate

Shape Circular colony having a diameter of about 1.5 mm after 24-hours incubation:

Rim:

Projection: Plain or hemispherical shape:

Positive; Gloss: Surface: Smooth:

Color: Creamy and semitransparent colony which forms no pink pigment;

(2) Characteristics of colony formed when incubated at 27°C in agar plate with dextrose and trypton Creamy and semitransparent colony with mucoid;

10 (3) Characteristics of colony formed when incubated at 27°C in agar plate with yeast extract and mannitol

Circular colony having a diameter of about 3 mm after 5-days incubation:

Color: Creamy and semitransparent colony with mucoid:

(4) Characteristics of colony formed when incubated at 27°C in agar plate with yeast extract, mannitol and congo

red Exhibiting neither a pale pink nor a substantial absorption of congo red;

(5) Growing at 27°C in agar plate with yeast extract, mannitol and 2% NaCl:

(6) Characteristics of colony formed when incubated at 27°C in slant nutrient agar

20 Growth: Satisfactory:

Shape: Thread-like; and

(7) Not liquefying gelatin when stab-cultured at 27°C in nutrient gelatin.

25 C. Physiological properties

15

35

40

45

50

- (1) Reduction of nitrate : Positive;
- (2) Denitrification reaction: Negative;
- (3) Methyl red test : Negative:
- 30 (4) VP-test : Negative;
 - (5) Formation of indole : Negative:
 - (6) Formation of hydrogen sulfide: Positive;
 - (7) Hydrolysis of starch: Negative;
 - (8) Utilization of citric acid: Positive:
 - (9) Utilization of inorganic nitrogen source: Utilizing ammonium salts and nitrates:

 - (10) Formation of pigment: Non; (11) Urease : Positive;
 - (12) Oxidase: Negative:
 - (13) Catalase : Positive;
 - (14) Growth conditions: Growing at a pH in the range of 5.5-9 0 and a temperature in the range of 4-35°C;
 - (15) Oxygen requirements : Aerobic;
 - (16) Utilization of carbon source and acid formation

Carbon source	Utilization	Acid formation
D-Glucose	+	+
D-Galactose	+	+
D-Fructose	+	+
L-Arabinose	+	+
D-Xylose	+	+
L-Rhamnose	+	+
Maltose	+	-
Sucrose	+	+
Lactose	+	-

(continued)

Carbon source	Utilization	Acid formation
Trehalose	+	-
Raffinose	+	+
Mannitol	+	-
Dextrin	+	-
Dulcitol	+	-

(17) Decarboxylase test on amino acid

Negative against L-lysine, L-arginine and L-omithine;

(18) Utilization of amino acid

5

10

15

20

25

45

50

55

Utilizing sodium L-glutamate, sodium L-asparate, L-histidine and L-proline;

(19) DNase: Negative;

(20) Formation of 3-ketolactose: Negative; and

(21) Mol% guanine (G) plus cytosine (C) of DNA: 61%.

The bacteriological properties were compared with those of known microorganisms with reference to Bergay's Manual of Systematic Bacteriology. Its delition (1984). As a result, it was revealed that the microorganism was identified as a microorganism of the genus Rhizobium. The microorganism is similar to those of the species Rhizobium melilot!, but it is distinguishable from them in that it utilizes maltose, lactose and mannitol but forms no acid, and in that it produces both an enzyme, which forms non-reducing seccharides having a trehalose structure when allowed oct or reducing partial starch hydrolysates, and a novel trehalose-releasing enzyme which specifically hydrolyzes the linkage between a trehalose molely and the remaining plycosyl moleiry in a non-reducing saccharide to lease the trehalose molely. No publication has reported such a microorganism having these properties.

The microorganism had been named "Rhizobium sp. M-11" by the present inventors and deposited on December 24, 1992, in Fermentation Research Institute, Agency of Industrial Science and Technology, Ibaraki, Japan. The deposition of the microorganism was accepted on the same day and has been maintained by the institute under the accession number of FERM BP-4130

In addition to the above-identified microorganism, other strains of the genus *Rhizobium* and their mutants can be sultably used in the invention as long as they produce the present trehalose-releasing enzyme.

The identification test of a microorganism of the genus Arthrobacter, i.e. Arthrobacter sp. Q36 according to the present invention gave the following results. The test was conducted in accordance with the method as described in "Biseibutsu-no-Bunru-i-to-Dotei" (Classification and Identification of Microorganisms), edited by Takeji Hasegawa, published by Japan Scientific Societies Press, Tokyo, Japan (1995). The results were as follows:

A. Morphology

(1) Characteristics of cells when incubated at 27 in nutrient agar

Usually exhibiting a rod form of 0.5-0.7x0.8-1.6µm; Existing single;

Exhibiting polymorphism;

Possessing no motility, flagellum and asporogenicity;

Non-acid fast;

Gram stain : Positive;

Capsule : Negative: and

(2) Characteristics of cells when incubated at 27°C in EYG agar

Exhibiting a rod-coccus cycle.

B. Cultural property

(1) Characteristics of colony formed when incubated at 27°C in nutrient agar plate

Shape Circular colony having a diameter of about 2-2.5 mm after 3-days incubation;

Rim: Entire:

Projection: Hemispherical shape;

Gloss: Moist gloss;

Surface: Smooth;

Color: Semitransparent and white or pale yellow;

(2) Characteristics of cells when slant-cultured at 27°C in nutrient agar plate

Growth rate : Satisfactory; and

10 Shape Thread-like;

5

15

30

40

45

50

55

(3) Characteristics of cells when slant-cultured at 27°C in agar plate containing yeast extract and peptone

Growth rate: Satisfactory;

Shape Thread-like; and

(4) Characteristics of cells when stub-cultured at 27°C in bouillon and gelatin Liquefying bouillon and gelatin.

20 C. Physiological properties

- (1) Reduction of nitrate : Positive:
- (2) Denitrification reaction : Negative;
- (3) Methyl red test : Negative;
- 25 (4) VP-test : Positive;
 - (5) Formation of indole : Negative;
 - (6) Formation of hydrogen sulfide : Positive;
 - (7) Hydrolysis of starch : Negative;
 - (8) Hydrolysis of cellulose: Negative:
 - (9) Utilization of citric acid : Positive;
 - (10) Utilization of inorganic nitrogen source Utilizing ammonium salts and nitrates:
 - (11) Formation of pigment : Negative;
 - (12) Urease : Positive:
 - (13) Oxidase : Negative;
 - (14) Catalase : Positive;
 - (15) Growth conditions: Growing at a pH in the range of 5-10 and a temperature in the range of 4-37°C;
 - (16) Oxygen requirements : Aerobic;
 - (17) Utilization of carbon source and acid formation

Utilization	Acid formation
+	-
+	-
+	-
+	-
+	-
+	-
+	-
+	-
+	-
+	-
+	-
+	-
	+ + + + + + + + + + + + + + + + + + + +

(continued)

Carbon source	Utilization	Acid formation
Dulcitol	+	-

(18) Utilization of amino acid

5

10

15

25

45

Utilizing sodium L-glutamate, sodium L-asparate, L-histidine and L-proline;

- (19) DNase : Positive:
- (20) Formation of 3-ketolactose : Negative;
 - (21) Major diamino acid of cell wall: Lysine: and
 - (22) Mol% guanine (G) plus cytosine (C) of DNA: 63%,

The bacteriological properties were compared with those of known microorganisms with reference to Bergey's Manual of Systematic Bacteriology Vol 2 (1986). As a result, I was revealed that the microorganism was identified as a microorganism of the genus Arthrobacter. The microorganism is characteristic in that it produces a non-reducing saccharide-forming anyme to form non-reducing saccharides having a trehables structure when allowed to act reducing partial starch hydrolysets, and a novel trehables-releasing enzyme which specifically hydrolyzes the lindage between a trehalose moiety and the remaining glycosyl moiety in a non-reducing saccharide to release the trehalose moiety. No publication has responded such a microorganism having these properties.

The microorganism had been named "Arthrobacter'sp. Q36" by the present inventors, and deposited on June 3, 1993, in National Institute of Bioscience and Human-Technology Agency of Industrial Science and Technology, Ibaraki, Japan. The deposition of the microorganism was accepted on the same day and has been maintained by the institute under the accession number of FERM BP-4316.

In addition to the above-mentioned microorganism, other strains of the genus Arthrobacter and their mutants can be suitably used in the invention as long as they produce the present trehabse-releasing enzyme which specifically hydrolyzes the linkage between a trehalose moiety and the remaining glycosyl moiety in non-reducing partial starch hydrolysaise having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher.

Other microorganisms can be used in the invention as long as they produce the present enzyme. For example, in addition to the above *Rhizobium* pp. M-11 (FERM BP-4130) and *Arthrobacter* sp. Q36 (FERM BP-4316), other hitherto known microorganisms such as those of the species *Brevibacterium helvolum* (ATCC 11822) and *Micrococcus roseus* (ATCC 186) can be favorably used in the invention.

Any nutrient culture medium can be used in the invention as long as such microorganisms can grow therein and produce the present trehalose-releasing enzyme: For example, synthetic- and natural-nutrient culture media can be arbitrarily used. Any carbon-containing substance can be used in the invention as a carbon source as long as it is utilized by the microorganisms: Examples of such a carbon source are saccharides such as glucose, fructose, lactose, sucrose, mannitol, sorbitol, molesses and parlial starch hydrobysates; and organis acids such as citic acid and succinic acid. The concentrations of these carbon sources in nutrient culture media are appropriately chosen. For example, in the case of using parlial starch hydrobysates, a preferable concentration is usually 20% or lower, more particularly, or lower, do s.b., in view of the growth and proliferation of the microorganisms. The nitrogen sources usable in the invention are, for example, inorganic nitrogen compounds such as ammonium salts and nitrates; and organic nitrogen-containing substances such as urea, com steep liquor, casein; peptone, yeast extract and beef extract. The inorganic ingredients usable in the invention are, for example, calcium salts, magnesium salts, potassium salts, sodium salts, phosphates and other salts of manganese, zinc, iron, copper, molybdenum and cobalt. If necessary, amino acids and vitamins can be favorably used.

The microorganisms usable in the invention are cultured under aerobic conditions at a temperature, usually a temperature in the range of 4-40°C, preferably, a temperature in the range of 20-35°C, and at a pH in the range of 4-10, preferably, a pH in the range of 5-9. The cultivation time suitably used in the invention is set to a time which longer than that required for the growth initiation of the microorganisms, preferably, 10-100 hours. The concentration of disactived oxygen (DC) in nutrient culture media is not specifically restricted, and usually a Do in the range of 200 ppm is satisfactorily used. The concentration of DO can be kept within the range by controlling the aeration rate, stirring nutrient culture media, supplementing oxygen to aeration, and increasing the inner pressure of fermenters. The culture can be carried out batchwise or in continuous manner.

After completion of the culture of microorganisms, the present enzyme is recovered Inasmuch as the activity of the present enzyme is found in both cells and cell-free supernatiants, they can be recovered and used as a crude enzyme. The resultant culture can be used intact as a crude enzyme. Conventional liquid-solid separation methods can be employed in the invention to remove cells from the culture. For example, methods to directly centrifuge the resultant cultures and those to filtrate them with precent filters or to separate cells by membrane filtration using plate filters or hollow fibers can be suitably used. Cell-free filtrates thus obtained can be used intact as an enzyme solution or they may be concentrated prior to their use. The concentration methods usable in the invention are for example, salting out using ammonium suitate, sedimentation using acetone and alcohol, and concentration using membranes such as plate filters and hollow fibers.

Coll-free filtrates and their concentrates can be subjected to conventional immobilization methods. Examples of conventional methods are conjugation methods using ion-exchangers, and covalent linkages and absorptions using resins and membranes, as well as inclusion methods using high-molecular weight substances. Cells separated from the resultant culture can be used as a crude enzyme without any further treatment or may be immobilized prior to their use. For example, the cells are immobilized by mixing them with sodium alignate, and dropping the resultant mixture in calcium chloride solution to gelatinize the drops into granules. The resultant granules can be fixed by using polyethylene imine or glutaraldehyde. Enzyme preparations extracted from cells can be used in the invention as a crude enzyme solution. Clear crude enzyme solutions containing the present enzyme can be obtained by extracting the present enzyme from cells, which were pretreated with ultrasonic, mechanical disruption using glass beads and alumina, front-press disruption, dec., and subjecting the resultant extracts to centrifugation or membrane filtration.

The crude enzyme solutions thus obtained can be used infract or purified by conventional methods prior to their use. For example, a purified enzyme preparation, which exhibits a single band on electrophoresis, can be prepared by dielzying a crude enzyme preparation, which were prepared by salting out the crude enzyme cultures with ammonium sulfate and concentrating the resultant, and successively purifying the delayzed solution on anion-exchange column chromatography using "DEAE" topopeatiffor, an anion exchange; hydrophobic column chromatography using "buyl-Toyopeariffor, a hydrophobic resin; and gel filtration chromatography using "Toyopeariffor and product of Tosoch Corporation, Toyo, Japan. Toyopeariffor, a resin for gel filtration, all of which are products of Tosoch Corporation, Toyop, Japan.

The present trehalose-releasing enzyme thus obtained has the following physicochemical properties:

(1) Action

15

25

30

40

Specifically hydrolyzing the linkage between a trehalose moiety and the remaining glycosyl moiety in a nonreducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher.

(2) Molecular weight

About 57,000-68,000 daltons on sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE);

(3) Isoelectric point (pI)

About 3.3-4.6 on isoelectrophoresis using ampholyte:

(4) Optimum temperature

About 35-45°C when incubated at pH 7.0 for 30 min;

(5) Optimum pH About 6.0-7.5 when incubated at 40°C for 30 min:

(6) Thermal stability Stable up to a temperature of about 30-45°C when incubated at pH 7.0 for 60 min; and

GIZDIE up

Stable at a pH of about 5 0-10.0 when incubated at 25°C for 16 hours.

The activity of the present trehalose-releasing enzyme is assayed as follows: One ml of an enzyme solution is added a 64 ml of 1.25 wV % maltoriosyltrehalose alias or-maltotertaceyli egiluccide in 50 mll phosphate buffer (pH 7.0), and the mixture solution is incubated at 40°C for 30 min. To the resultant reaction mixture is added a copper solution for the Somogyli reaction to suspend the enzymatic reaction, followed by the determination of the reducing power on the Somogyli-Nelson's method. As a control, an enzyme solution, which were preheated at 100°C for 10 min to inactivate the enzyme, is assayed similarly as above. One unit activity of the present enzyme is defined as the amount of enzyme which increases the reducing power of that of one jumole of glucose per minute when assayed with the above-methored assay.

Any substance can be used as a substrate for the present enzyme as long as it is a non-reducing seacharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher. Examples of such a substrate are glycosyltrehaloses such as glucosyltrehalose, malicotyriehaloses, malicotyriehaloses and maltopentacsyltrehaloses which are prepared by allowing a non-reducing seacharide-forming enzyme to a con mallotoriese. malloterlaces, malloterlaces, mallohoxaces and maltohepatoses in a ddition to these substrates, relatively-low reducing partial starch hydrolysates, having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher, which can be prepared by partially hydrolyzing armylaceous substances such as faterd, armylopedin and armylose with armylases or acids can be suitably used in the invention.

Examples of such amylases which partially hydrolyzing starch are α-amylase, maltopentaose-forming amylase and maltohexaose-forming amylase as disclosed in *Handbook of Amylases and Related Enzymes*, published by Per-

gamon Press, Tokyo, Japan (1988). These amylases can be used in combination with debranching enzymes such as pullulanase and isoamylase.

The concentration of the reducing partial starch hydrolysates used as a substrate in the invention is not specifically restricted. The enzymatic reaction according to the present invention proceeds even in a solution containing of 10% or 50% of a substrate to form trehalose. Suspensions containing insoluble substrates can be used in the invention. The reaction temperature usable in the present enzymatic reaction can be set to a temperature at which the present enzymatic is not inactivated, it, a a temperature up to about 55°C, preferably, a temperature in the range of 40°-55°C. The reaction pt usable in the present enzymatic reaction is set to a pt in the range of 5°-10, preferably, a pt in the range of about 6°-8. The reaction inter usable in the present enzymatic reaction is adequately chosen dependently on the conditions of the enzymatic reaction, and, usually it is in the range of about 0.1°-100 hours when the present enzyme is used in a mornout of about 0.1°-100 intellig substrate, d. s.b.

As regards the yield of trehalose from material substrates, specifically, in the case of preparing trehalose from nonreducing partial starch hydrolysates with a relatively-low DE value, i.e. those with a relatively-high degree of glucose polymerization, the present preparation of trehalose has the advantage of that it increases the yield of trehalose more than that attained by a preparation as discobsed in Japanese Patent Application No.382, 131/92 wherein a non-reducing saccharied-forming enzyme and glucoamy/lase are used in combination. The present preparation, wherein a non-reducing saccharide-forming enzyme and the present trehalose-releasing enzyme are used in combination, forms trehalose in a highly yield of about 60% or higher, while the preparation of the Japanese Patent Application forms trehalose only in a vield of about 60%.

The enzymatic mechanism of the present invention is as follows: A reducing partial starch hydrolysate with a relatively-high degree of glucose polymerization is first converted by a non-reducing saccharide-forming enzyme into one mole of a non-reducing saccharide having a trehalose structure as an end unit, then the resultant non-reducing saccharide is hydrolyzed by the present trehalose-releasing enzyme into one mole of a reducing partial starch hydrolyzed by the present trehalose-releasing enzyme into one mole of a reducing partial starch hydrolyzed by 2 in the case of that the newly formed reducing partial starch hydrolyzed has a degree of glucose polymerization of 10 or higher, it can be further converted into a non-reducing saccharide having a trehalose structure as an end unit, and then converted into one mole of trehalose and a partial starch hydrolyzed by the trehalose-releasing enzyme. Accordingly, repeated enzymatic reactions of the altoresate in on-reducing saccharide-forming onzyme and trehalose-releasing enzyme an form from one mole of a non-reducing partial starch hydrolyzeate with a degree of glucose polymerization of lower than that of the material partial starch hydrolystate by a number of 2-fold higher than that of the formed trehalose-releasing enzyme.

25

45

55

In the present preparation, a non-reducing saccharide-forming enzyme and the present trehalose-releasing enzyme can be simultaneously allowed to act on non-reducing partial starch hydrolysates having a degree of glucose polymerization of 3 or higher, or those two enzymes can be successively allowed to act on non-reducing partial starch hydrolysates in this order. In order to more increase the trehalose yield, the resultant reaction mixture can be further subjected to the action of glucoarnylase.

The reaction mixtures thus obtained are in usual manner subjected to filtration and centrifugation to remove insoluble substances, and the resultant solutions are decolored with an activated charcoal, desalted with in-exchanges in H-and OH-form, and concentrated into syrupy products. The syrupy products can be arbitrarily dried into powdery nordurs.

If necessary, the powdery products are readily processed into high-purity trehalose products by purifying them with one or more methods, for example, fractionations on ion-exchange column chromatography, column chromatography using an activated charcoal or a silica gel; separations using organic solvents such as alcohol and acetone; and alkaline treatments to decompose and remove the remaining reducing seacharides.

If necessary, the saccharide products containing the trefaclose according to the invention can be hydrolyzed by campiase, Busanylase, glucanylase, explusorations and/or trefalse, or subjected to a saccharide-transfer reaction by using cyclomaltodextrin glucanotransferase and/or glucosyltransferase to control their sweetness and reducing power as well as to reduce their viscosity. Furthermore, the saccharide products can be arbitrarily hydrogenated to conveol them into sugar alcohols to eliminate their reducing power. From the resultant products glucose can be removed by using aforesaid purification methods such as ion-exchange column chromatography to prepare high trefalsice content fractions. The fractions thus obtained can be readily purified and concentrated into syrupy products, and, if necessary the syrupy products can be further concentrated into supersaturated solutions and crystallized to obtain hydrous crystalline trehalose or rahydrous crystalline trehalose or rahydrous crystalline trehalose.

The ion-exchange column chromatographic techniques usable in the invention include, for example, those which employ a strong-acid caltion-exchange resin as disclosed in Japanese Patent Laid-Open Nos 23,799/83 and 72,599/83. By using the techniques, concomitant saccharides contained in crude trehalose products can be readily removed to obtain high trehalose content products. In this case, any one of fixed-bed, moving bed and semi-moving methods can be arbitrarily employed.

15

25

35

40

45

50

55

In order to prepare hydrous crystalline trehalose, an about 65-90% trehalose solution is placed in a crystallizer, and gradually cooled while stirring in the presence of 0.1-20% seed crystal at a temperature of 95°C or lower, preferably, a temperature in the range of 10-90°C. to obtain a massecuite containing hydrous crystalline trehalose. Conventional methods such as separation, block pulverization, fluidized-bed granulation and spray drying can be employed in the invention to propare from the masseculte hydrous crystalline trehalose or crystalline saccharides containin at I

In the case of separation, massecuites are usually subjected to a basket-type centrifuge to separate hydrous crystalline trehalose from the mother liquor, and, if necessary the hydrous crystalline trehalose is washed by spraying it with a small amount of cold water to facilitate the preparation of hydrous crystalline trehalose with an increased purity. In the case of spray drying, crystalline saccharides with no or substantially free of hygroscopicity are readily prepared by spraying massecuites, having a concentration of 60-85%, d.s.b., and a crystallinity of about 20-60%, d.s.b., from a nozzle by a high-pressure pump; drying the resultants with an about 60-100°C hot air which does not melt the resultant crystalline powders, and aging the resultant powders for about 1-20 hours while blowing thereto an about 30-60°C hot air. In the case of block pulverization, crystalline saccharides with no or substantially free of hygroscopicity are readily prepared by allowing massecuites, having a moisture content of about 10-25% and a crystallinity of about 10-60%, d. s.b., to stand for several hours to 3 days or so in order to crystallize and solidify the whole contents into blocks, pulverizing or cutting the resultant blocks, and drying the resultants. Although anhydrous crystalline trehalose can be prepared by drying hydrous crystalline trehalose to convert it into anhydrous form, it is generally prepared by providing a high trehalose content solution with a moisture content less than 10%, placing the solution in a crystallizer, keeping the solution in the presence of a seed crystal at a temperature in the range of 50-160°C, preferably, a temperature in the range of 80-140°C under stirring conditions to obtain a massecuite containing anhydrous crystalline trehalose, crystallizing it, and pulverizing the resultant anhydrous crystalline trehalose under dryness and relatively-high temperature conditions by conventional methods such as block pulverization, fluidized-bed granulation and spray drying.

The present trehalose thus obtained is stable and substantially free of reducing power, and can be mixed and processed with other materials, specifically, amine acids and amine acid-containing substances such as oligopeptitides and proteins without a fear of causing unsatisfactory browning and smell as well as deterioration of the materials. Trehalose per se has a satisfactorily-high quality and sweetness. Short brehalose is readily hydrolyzed by trehalases into glucose units, it is assimilated, absorbed and utilized by living bodies as a caloric source when orally administer. Furthermore, trehalose is not substantially fermented by dental carries-inducing microorganisms, and this renders it useful as a sweetner substantially free of inducing dental carries.

The present trehalose can be prepared into agents, for example, nutritional agents for transfusion and intubation feeding, which are arbitrarily administrable to living bodies and readily metabolized and utilized by the living bodies without a fear of causing toxicity and side effects. Thus, these products can be advantageously used as an energy-supplementing agent for living bodies.

Trehalose is a stable sweetener, and, especially crystalline trehalose is arbitrarily used as a sugar coating agent for tablets when used in combination with a binder such as pullulan, hydroxyethyl starch or poty/inylpyrroidone. In addition, trehalose has properties such as osmotic pressure-controlling ability, filler-imparting ability, olioss-imparting ability, moisture-retaining ability, viscosity-imparting ability, substantial no fermentability, ability to prevent retrogradation of gelatinized starch, and ability to prevent or substitutions of the saccharides.

Thus, the present trahalose and saccharide composition containing the same can be arbitrarily used as a sweetener, taste-improving agent, quality-improving agent, stabilizer and filler in a variety of compositions such as food products, cigarettes, tobaccos, feeds, cosmetics and pharmacouticals.

The present trehalose and saccharide compositions containing the same can be used intact as a seasoning for sweetening. If necessary, they can be used together with adequate amounts of one or more other sweeteners, for example, powdered syrup, glucose, fructose, maltose, sucrose, isomerized sugar, honey, maple sugar, erythritol, sorb-tiol, matittol, lactitol, dihydrocharone, stevioside, c-glycosyl stevioside, rebaudioside, glycyrritizin, L-aspartyl L-phenylatanine methyl ester, saccharin, glycine and alanine, and/or a filler such as dextrin, starch and lactose.

The present trehalose and saccharide compositions containing the same in the form of a powder or a crystal can be used intact, or, if necessary they can be mixed with an excipient, filler, diluent and binder and formed into granules, spheres, shot-rods, plates, cubes and tablets, prior to their use.

The present trehalose and seacharide compositions containing the same well harmonize with other materials having sour, acid, sally, biller, satingent- and delicious-tastes, and have a relatively-high acid tolerance and heat resistance. Thus, they can be favorably used in food products in general as a sweetener, taste-improving agent and quality-improving agent.

The present trehalose and saccharide compositions containing the same can be used in seasonings such as an amino acid, peptide, soy sauce, powdered soy sauce, "miso", "furnnatu-miso" (a powdered miso), "moromi" (a refined sake), "hishio" (a refined soy sauce), "furikako" (a seasoned fish meal), mayonnaise, dressing, vinegar, "sanbai-zu" (a sauce of sugar, soy sauce and vinegar), "furnnatu-uschi-su" (powdered vinegar for sushi), "chuka-no-moto" (an instant

mix for Chinese dish), "tentsuyu" (a sauce for Japanese deep-fat fried food), "mentsuyu" (a sauce for Japanese vermicelli), sauce, catsup, "yakiniku-no-tare" (a sauce for Japanese grilled meat), curry roux, instant stew mix, instant soup mix, "dashi-no-moto" (an instant stock mix), nucleic acid condiments, mixed seasoning, "mirin" (a sweet sake), "shin-mirin" (a synthetic mirin), table sugar and coffee sugar.

The present trehalose and saccharide compositions containing the same can be also used freely for sweetening "wagashi" (Japanese cakes) such as "senbei" (a rice cracker), "arare-mochi" (a rice-cake cube), "okoshi" (a millet-andrice cake), "mochi" (a rice paste), "manju" (a bun with a bean-jam), "uiro" (a sweet rice jelly), "an" (a bean jam), "yokan" (a sweet jelly of beans), "mizu-vokan" (a soft adzuki-bean jelly), "kingvoku" (a kind of yokan), jelly, pao de Castella and "amedama" (a Japanese toffee); confectioneries such as bun, biscuit, cracker, cookie, pie, pudding, butter cream, custard cream, cream puff, waffle, sponge cake, doughnut, chocolate, chewing gum, caramel and candy; frozen desserts such as ice cream and sherbet; syrups such as "kajitsu-no-syrup-zuke" (a preserved fruit) and "korimitsu" (a sugar syrup for shaved ice); pastes such as flour paste, peanut paste, fruit paste and spread; processed fruits and vegetables such as jam, marmalade, "syrup-zuke" (fruit pickles) and "toka" (conserves), pickles and pickled products such as "fukujin-zuke" (red colored radish pickles), "bettara-zuke" (a kind of whole fresh radish pickles), "senmai-zuke" (a kind of sliced fresh radish pickles) and "rakkyo-zuke" (pickled shallots); premixes for pickles and pickled products such as "takuan-zuke-no-moto" (a premix for pickled radish) and "hakusai-zuke-no-moto" (a premix for fresh white rape pickles); meat products such as ham and sausage; products of fish meat such as fish ham, fish sausage, "kamaboko" (a steamed fish paste), "chikuwa" (a kınd of fish paste) and "tenpura" (a Japanese deep-fat fried fish paste); "chinmi" (relish) such as "uni-no-shiokara" (salted guts of sea urchin), "ika-no-shiokara" (salted guts of squid), "sukonbu" (processed tangle), "saki-surume" (dried squid strips) and "fugu-no-mirin-boshi" (a dried mirin-seasoned swellfish); "tsukudani" (foods boiled down in soy sauce) such as those of laver, edible wild plants, dried squid, fish and shellfish; daily dishes such as "nimame" (cooked beans), potato salad and "konbu-maki" (a tangle roll); milk products; canned and bottled products such as those of meat, fish meat, fruit and vegetable; alcoholic beverages such as synthetic sake, wine and liquors; soft drinks such as coffee, tea, cocoa, juice, carbonated beverage, sour milk beverage and beverage containing a lactic acid bacterium; instant food products such as instant pudding mix, instant hot cake mix and "sokuseki-shiruco" (an instant mix of adzuki-bean soup with rice cake) and instant soup mix; and beverages such as baby foods, foods for therapy, and beverages supplemented with nutrition; as well as for improving the tastes and qualities of the aforementioned food-products.

15

25

30

35

45

50

55

The present trehalose and saccharide compositions containing the same can be also used in feeds and pet foods for animals such as domestic animals, poultry, honey bees, silk worms and fishes in order to improve their taste preferences. The trehalose and saccharide compositions can be arbitrarily used as a sweetener, taste-improving agent, quality-improving agent and stabilizer in other products in paste and liquid form such as a tobacco, cigarette, dentifrice, lipstick, rouge, lip cream, internal medicine, tablet, troche, cod liver oil in the form of a drop, cachou, oral refrigerant, gardle, cosmetic and pharmacountical.

The present trehalose and saccharide compositions containing the same can be used as a quality-improving agent and stabilizer for biologically active substances susceptible to loss of their effective ingredients and activities, as well as in health foods and pharmaceutical compositions containing biologically active substances. Examples of such a biologically active substance are lymphokines such as α -, β -and γ -interferons, tumor necrosis factor- α (TNF- α), tumor necrosis factor-β (TNF-β), macrophage migration inhibitory factor, colony-stimulating factor, transfer factor and interleukin 2 (IL-2); hormones such as insulin, growth hormone, prolactin, erythropoietin and follicle-stimulating hormone; biological preparations such as BCG vaccine, Japanese encephalitis vaccine, measles vaccine, live polio vaccine, smallpox vaccine, tetanus toxoid, Trimeresurus antitoxin and human immunoglobulin; antibiotics such as penicillin, erythromycin, chloramphenicol, tetracycline, streptomycin and kanamycin sulfate; vitamins such as thiamine. riboflavin, L-ascorbic acid, cod liver oil, carotenoid, ergosterol and tocopherol; enzymes such as lipase, elastase, urokinase, protease, B-amylase, isoamylase, glucanase and lactase; extracts such as ginseng extract, snapping turtle extract, chlorella extract, aloe extract and propolis extract; viable microorganisms such as viruses, lactic acid bacteria and yeasts; and other biologically active substances such as royal jelly. By using the present trehalose and saccharide compositions containing the same, the aforementioned biologically active substances are readily prepared into health foods and pharmaceutical compositions with a satisfactorily-high stability and quality without a fear of losing or inactivating their effective ingredients and activities

As described above, the methods to incorporate the present trehalose and saccharide compositions containing the same into the adorementioned substances and compositions include conventional methods, for example, mixing, kneading, dissolving, melting, soaking, permeating, sprinkling, applying, coating, spraying, injecting, crystallizing and soliditying. The trehalose and saccharide compositions containing the same are usually incorporated into the aforementoned substances and compositions in a manuout of 0.1% or higher, fortedpily, one % or higher, d.s.b.

The following experiments explain the production and purification of the present trehalose-releasing enzyme derived from microorganisms of the species *Rhizobium* sp. M-11 and *Arthrobacter* sp. Q35, as well as hitherto known microorganisms

Experiment 1

Production of trehalose-releasing enzyme by Rhizobium sp. M-11

A liquid nutrient culture medium, consisting of 2 0 w/v % "PINE-DEX #4", a starch product of Matsusian Chemical Ind., Co., Ltd., Kyoto, Japan, 0.5 w/v % peptone, 0.1 w/v % yeast extract, 0.1 w/v % disodium hydrogenphosphate, 0.1 w/v % potassium hydrogenphosphate and water, was adjusted to pH 7.0 About 100 m laliquots of the liquid nutrient culture medium were placed in 500-ml Eflenmeyer flasks, autoclaved at 120°C for 20 min to effect sterilization, cooled, inoculated with a seed culture of *Phizobium* sp. M-11 (FERIM BP-4130), and incubated at 27°C for 24 hours under stirring conditions of 130rpm. The resultant cultures were pooled and used as a seed culture.

About 20 L of a fresh preparation of the same liquid nutrient culture medium as used in the above culture was placed in a 30-L lementer, sterilized, cooled to 27°C, incoulated with one w/v % of the seed culture, and incubated for about 72 hours under striring and aerobic conditions at 27°C and a 04 to 6.0-8.0.

The activities of a non-reducing saccharide-forming enzyme and the present trehalose-releasing enzyme accumulated in the culture were respectively about 1.5 units/ml and about 2 units/ml. A portion of the culture was separated by centrifugation into cells and a supernatant, and the cells were suspended in 50 mM phosphate bufler (p1.7.0) to give the same volume of the portion, followed by assaying the enzyme activity of the cell suspension and the supernatant. The activities of the non-reducing saccharide-forming enzyme and the present trehalose-releasing enzyme in the cell suspension were respectively about 0.6 units/ml and about 0.9 units/ml, and the culture supernatant contained about 0.9 units/ml of the non-reducing saccharide-forming enzyme and about 1.2 units/ml of the present trehalose-releasing enzyme.

The assay of the non-reducing saccharide-forming enzyme is as follows: One mI of an enzyme solution is addid to 4 mI of 1.28 w. % maltopentace in 50 mM phosphate buffer (pH 7.0), and the mixture solution is incubated at 4PCC for 10 min and heated at 100°C for 10 min to suspend the enzymatic reaction. The resultant reaction mixture is precisely diluted with a buffer by 10 times, and the reducing power was determined on the Somogyi-Nelson's method. As a control, an enzyme solution, proheated at 100°C for 10 min to inactivate the enzyme, is assayed similarly as above. One unit activity of the non-reducing saccharide-forming enzyme is defined as the amount of enzyme which eliminates the reducing power of that of one purpole or implicentaces per mixture when assayed with the above-mentioned assay.

Experiment 2

15

35

45

Purification of enzyme

An about 18 L of a culture obtained by the method in Experiment 1 was treated to disrupt cells with "MINI-RABO", a supper high-pressure cell disrupting apparative commercialized by Dainippon Pharmaceutical Co., Ltd. Tokyo, Janan. The resultant suspension was centrifuged at 10,000:pm for 30 min to obtain an about 16 L supernatant. Ammonium sulfate was added to the supernatant and dissolved therein to give a saturation degree of 0.2, and the resultant solution was allowed to stand at 4°C for one hour, and centrifuged at 10,000:pm for 30 min to obtain a supernatant.

Ammonium sulfate was actided to the resultant supernatant and dissolved therein to give a saturation degree of 0.6, and the resultant solution was centrifuged at 10.000rpm for 30 min, followed by recovering a precipitate and dissolving it in 10 mM phosphate buffer (pH 7.0). The solution thus obtained was dialyzed against a fresh preparation of the same phosphate buffer for 24 hours, and centrifuged at 10.000rpm for 30 min to remove insolution substances. Three hundred and sixty mil of the resultant dialyzingd solution was divided into 2 portions which were then separately subjected to column chromatography using a column packed with 300 mll of "DEAE-Toyopearl®", an ion-exchanger commercialized by Tosoh Corporation, Tokyo, Japan.

The objective non-reducing saccharide-forming enzyme and trehalose-releasing enzyme were adsorbed on the ion-exchanger, and eluted separately from the column with a fresh preparation of the same phosphate buller supplemented with sait at different sait concentrations. The elution pattern of the column or the column chromatogram was as shown in FIG.1. The non-reducing saccharide-forming enzyme was eluted from the column at a sait concentration of about 0.2 M while the trehalose-releasing enzyme was eluted from the column at a sait concentration of about 0.3 M. The fractions containing either of the objective enzymes were separately pooled and purified.

The pooled fractions containing the non-reducing saccharide-forming enzyme were dialyzed against a fresh preparation of the same phosphate buffer supplemented with 2 M ammonium suttlest. The dialyzed solution was centrifuged at 10 000rpm for 30 min to remove insoluble substances, and the resultant supernatant was subjected to hydrophobic column chromatography using a column packed with 300 ml of "Sutyl-Toyopeari@550"; a hydrophobic pel commercialized by ToseOh Corporation, Tokyo, Japan. The enzyme adsorbed on the gel was eluted from the column with a line gradient buffer ranging from 2 M to 0 M, followed by recovering fractions with the enzyme activity. The resultant fractions were subjected to gel filtration chromatography using a column packed with 300 ml of Toyopeari@HV-55"; a resin for

gel chromatography commercialized by Tosoh Corporation, Tokyo, Japan, followed by recovering fractions with the enzyme activity.

By using the pooled fractions with a trehalose-releasing enzyme activity eluted from the column of "DEAE-Toyopearl®", the fractions were treated similarly as in the purification steps used in the preparation of the non-reducing saccharide-forming enzyme in such a manner that they were dialyzed against a buffer containing 2 M ammonium sulfate, and successively subjected to hydrophobic column chromatography and gel filtration chromatography.

The total enzyme activity, specific activity and yield of the non-reducing saccharide-forming enzyme in each purification step are as shown in Table 1, while those of the trehalose-releasing enzyme are as shown in Table 2.

		Table 1		
	Purification step	Total enzyme*activity (units)	Specific activity (units/mg protein)	Yield (%)
	Material culture	28,500	-	100
	Supernatant after cell disruption	22,900	0.12	80
	Dialyzed solution after salting out	21,100	0.43	74
	Eluate from ion- exchange column	15,200	6.2	53
	Eluate from hydrophobic column	7,950	101	28
;	Eluate after gel filtration column	5,980	197	21

Note: The symbol "" means a non-reducing saccharide-forming enzyme.

T-11- 0

30		Table 2		
	Purification step	Total enzyme** activity (units)	Specific activity (units/mg protein)	Yield (%)
	Material culture	37,400	-	100
35	Supernatant after cell disruption	31,500	0.17	84
	Dialyzed solution after salting out	29,200	0.60	78
40	Eluate from ion- exchange column	25,400	5.3	68
	Eluate from hydrophobic column	18,700	98.5	50
45	Eluate from gel filtration column	11,600	240	31

Note: The symbol "**" means the present trehalose-releasing enzyme.

The purified enzyme preparations, obtained as an eluate from cell filtration column in Tables 1 and 2, were examined their purity on electrophoresis using 7.5% polyacrylamide gel. As a result, each enzyme preparation was observed as a single protein band, and this meant that it was an electrophoretically-homogeneous preparation with a relatively-high purity.

Experiment 3

55

10

15

20

25

Property of trehalose-releasing enzyme

A portion of a purified trehalose-releasing enzyme preparation, obtained by the method in Experiment 2, was

subjected to electrophoresis using a gel containing 10% sodium dodecylsulfate polyacrylamide, and determined its molecular weight to be about 57,000-68,000 deltons by comparing it with marker proteins commercialized by Japan Bio-Rad Laboratories. Tolivo, Japan

Another portion of the purified enzyme preparation was subjected to iscelectrophoresis using polyacylamide gel containing 2 w/v % "AMPHOLINE", an ampholyte commercialized by Pharmacia LKB Biotechnology AB, Uppsala, Sweden. The resultant gel was sliced into pieces, followed by measuring their pHs and resulting in a pl of the enzyme being about 3.94.3.

Effects of temperature and pH on the activity of the present enzyme were studied in accordance with the assay for the enzyme activity. These results were respectively shown in FIG. 2 (effect of temperature) and FIG. 3 (effect of pH). The optimum temperature of the enzyme was about 45°C when incubated at pH 7.0 for 30 min, and the optimum pH was about 6.0-7.5 when incubated at 40°C for 30 min. The thermal stability of the enzyme was determined by incubating it in 50 mM phosphate buffers (pH 7.0) for 60 min at different temperatures, cooling the buffers the studies with cold water, and determining the remaining enzyme activity in each buffer. The pH stability of the enzyme was determined by incubating it in 50 mM phosphate buffers having different pHs at 25°C for 16 hours, adjusting the buffers to pH 7, and assaying the remaining enzyme activity in each buffer. The results of the thermal- and pH-stabilities of the enzyme were respectively shown in FIG.s 4 and 5. The enzyme was stable up to a temperature of about 40°C and at a pH of about 5-10.

Experiment 4

20

25

40

50

55

Preparation of trehalose from non-reducing saccharides having a trehalose structure as an end unit and a degree of glucose polymerization of 3 or higher

Non-reducing saccharides, having a trehalose structure as an end unit and a degree of glucose polymerization of 3 or higher used as a substrate, were prepared according to the method as described in Japanese Patent Application No.362,131/92. To an aqueous solution containing 20% maltotriose, maltotetraose, maltopentaose, maltohexaose or maltoheptaose as a substrate was added 2 units/g substrate, d.s.b., of a purified enzyme preparation obtained by the method in Experiment 2, and the resultant mixture was subjected to an enzymatic reaction at 40°C and pH 7.0 for 48 hours. The reaction mixture was heated to inactivate the remaining enzyme, filtered, decolored, desalted and concentrated to obtain a concentrated saccharide solution which was then column chromatographed by using "XT-1016 (Na+form, polymerization degree of 4%)*, an ion-exchanger commercialized by Tokyo Organic Chemical Industries, Ltd., Tokyo, Japan. In the column chromatography, the ion-exchanger was packed in 3-jacketed stainless-steel columns, having an inner diameter of 2.0 cm and a length of one m, which were then cascaded in series, heated to give the inner column temperature of 55°C, applied with 5 v/v % of the concentrated saccharide solution against the resin while keeping at 55°C, and fed with 55°C hot water at SV (space velocity) of 0.13 to obtain high-purity non-reducing saccharides having a trehalose structure as an end unit and having a degree of polymerization of 3 or higher. Among the resultant preparations, a glucosyltrehalose preparation contained glucosyltrehalose with a purity of 97.6%, d.s.b., and the purities of maltosyltrehalose, maltotriosyltrehalose, maltotetraosyl-trehalose and maltopentaosyltrehalose in their high-purity preparations were respectively 98.6%, 99.6%, 98.3% and 98.1%, d.s.b.

An acueous solution containing 20%, d.s.b., of each one of the above 5 non-reducing saccharide preparations, namely glocosythenaloes preparations, ease prepared, followed by mixing it with 2 uniting substate, d.s.b. the purified trehalose-releasing enzyme obtained in Experiment 2, and subjecting the resultant solution to an enzymatic reaction at 40°C and PH 7.0 for 48 hours. The resultant each reaction mixture was desalted, and analyzed for its composition on high-performance liquid chromatography (HPLC) using "WAKOBEADS WB-T-330 column", a column of Wako Pure Chemical Industries Ltd., Tokyo, Japan. As a control, a fresh preparation of the same trehalose-releasing enzyme was allowed to act on mallooligosecharides such as mallotriose, malloterases, mallooperatoses, mallothexaces and malloheptaces, and the resultant each reaction mixture was analyzed for its composition on HPLC. The results were as shown in Table 3.

Substrate	Product	Elution time on HPLC (min)	Percentage (%)
	Trehalose	27.4	17.5
Glucosyltrehalose	Glucose	33.8	6.5
	Glucosyltrehalose	23.3	76.0
	Trehalose	27.4	44.3
Maltosyltrehalose	Maltose	28.7	44.4
	Maltosyltrehalose	21.6	11.3
	Trehalose	27.4	39.5
Maltotriosyltrehalose	Maltotriose	25.9	60.0
	Maltotriosyltrehalose	19.7	0.5
	Trehalose	27.4	34.2
Maltotetraosyltrehalose	Maltotetraose	24.1	65.5
	Maltotetraosvltrehalose	e 18.7	6.0

_
v
0
2
5
Ξ.
č
ö
Ö
-

Substrate	Product E	Elution time on HPLC (min)	Percentage (%)	
	Trehalose	27.4	29.1	
Maltopentaosyltrehalose	Maltopentaose	22.6	70.6	
	Maltopentaosyltrehalose 17.8	e 17.8	0.3	
Maltotriose	Maltotriose	25.9	100	
Maltotetraose	Maltotetraose	24.1	100	
Maltopentaose	Maltopentaose	22.6	100	
Maltchexaose	Maltohexaose	21.8	100	
Maltcheptaose	Mal toheptaose	21.0	100	

The results in Table 3 evidently show that:

- Trehalose-releasing enzyme according to the present invention specifically hydrolyzes the linkage between a
 trehalose moiety and a glycosyl moiety in a non-reducing saccharide having a trehalose structure as an end unit
 and having a degree of glucose polymerization of 3 or higher to form trehalose and a non-reducing saccharide
 having a degree of glucose polymerization of one or more; and
- 2. Maltooligosaccharides are not hydrolyzed by the present trehalose-releasing enzyme.

From these results, it is confirmed that the trehalose-releasing enzyme according to the present invention is a novel enzyme which has a mechanism of specifically hydrolyzing the linkage between a trehalose moiety and the remaining glycosyl moiety in a non-reducing saccharide having a trehalose structure as an end unit and having a degree of dlucose polymerization of 3 or higher to release trehalose from the non-reducing saccharide.

In order to purify trehalose in each reaction mixture, the reaction mixture was subjected to column chromatography using a column packed with "XT-1016 (Na*-form)", an alkaline-metal strong-acid cation exchange resin commercialized by Tokyo Organic Chemical Industries, Ltd., Tokyo, Japan, followed by recovering fractions containing 97% or higher of trehalose. The fractions were pooled and concentrated into an about 65% solution, and the concentrate was allowed to stand at 25°C for 2 days to crystallize hydrous crystalline trehalose, followed by separating and drying it in vacuo to obtain a high-purity trehalose preparation with a purity of 99% or higher, d.s.b. The yields of trehalose from glucosyltrehalose, maltosyltrehalose, maltotriosyltrehalose, maltotetraosyltrehalose and maltopentaosyltrehalose used as a substrate were respectively 9.5%, 14.9%, 16.0%, 18.5%, and 17.7%, d.s.b. The high-purity trehalose preparations and a commercially available trehalose specimen as a standard were studied on their melting point, heat of fusion, specific rotation, infrared absorption spectrum, powdery x-ray diffraction pattern, and readiness of hydrolysis by a trehalase specimen derived from pig kidney, commercialized by Sigma Chemical Co., St. Louise, USA. As a result, every trehalose preparation showed a melting point of 97.0±0.5°C, a heat of fusion of 57.8±1.2 kJ/mole and a specific rotation of +182±1.1°, and these values well corresponded with those of the standard trehalose specimen, while the infrared absorption spectra and powdery x-ray diffraction patterns of the trehalose preparations also well corresponded with those of the standard trehalose specimen. Similarly as the standard trehalose specimen, the trehalose preparations were decomposed into glucose units.

As evident from these results, it was confirmed that a non-reducing saccharide, which was formed by allowing the present trehalose-releasing enzyme to act on non-reducing saccharides having a trehalose structure as an end unit and having a degree of ollucose polymerization of 3 or higher, was trehalose.

Experiment 5

5

15

25

Preparation of trehalose from non-reducing partial starch hydrolysates

A suspension containing 5% waxy corn starch was gelatinized by heating, adjusted to pH 4.5, heated to 50°C, mixed with 4.000 units/g starch, d.s.b., of an isoamylase specimen commercialized by Hayashibara Biochemical Laboratories Inc., Okayama, Japan, and subjected to an enzymatic reaction for 20 hours. The reaction mixture was autoclaved at 120°C for 10 min, cooled to 60°C, and subjected to gel filtration column chromatography using a column packed with 750 ml of "Toyopeario" 50°S gel", commercialized by Tosch Corporation, Tokyo, Japan, to obtain reducing partial starch hydrolysates having a degree of glucose polymerization of 35-10.

Either of the reducing partial starch hydrolysates thus obtained or mallotriose having a degree of glucose polymerization of 3 as a substrate was dissolved in 10 mM phosphate buffer (pH 7.0) into a one % solution which was then mixed with 4 units/g substrate, d.s.b., of a purified non-reducing saccharide-forming enzyme and a purified trehalose-releasing enzyme prepared by the method in Experiment 2, and subjected to an enzymatic reaction at 40°C for 24 hours. After completion of the enzymatic reaction, a portion of the resultant each reaction mixture was desaited and analyzed on HPLC to identify its composition.

The remaining each reaction mixture was heated to 50°C, adjusted to pH 4.5, admixed with 50 units/g substrate, d.s.b, of a glucoamylase specimen commercialized by Selikagaku-Kogyo Co, Ltd, Tokyo, Japan, and subjected to an enzymatic reaction for 24 hours. Similarly as above, a portion of the resultant each reaction mixture was desalted and analyzed on HPLC to analyze its composition. The results were as shown in Table 4.

55

Degree of glucose polymerization of	Reaction product	Composition	tion (%)
reducing partial starch hydrolysate		P.	'n
	Trehalose	80.8	83.5
34.1	Reducing oligosaccharides Glycosyltrehalose	4.6	000
	Trehalose	79.7	82.5
26.2	Reducing oligosaccharides Glycosyltrehalose	15.3	200
	Trehalose	77.7	80.7
18.1	Reducing oligosaccharides Slycosyltrehalose	17.0	200
	Trehalose	75.0	78.5
15.2	Reducing oligosaccharides Glycosyltrehalose	18.6	00
	Trebalose	65.1	70.1
10.0	Reducing oligosaccharides Glycosyltrehalose	27.6	,00 ,00
(Trehalose	4.2	20.8
3 (Maltotriose)	Glucose Maltotriose Glucosyltrebalose	6 2.1 28.0 28.0	20.0

5

10

15

20

25

30

35

40

45

50

55

: The symbol "*" means a composition after enzymatic reaction of a non-reducing saccharide-forming enzyme and the present trehalose-releasing enzyme, and the In the Table, the wording "Glycosyltrehalose" means non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose symbol "**" means a composition after enzymatic reaction of glucoamylase. polymerization degree of 3 or higher. Note

As is shown in Table 4, in the case of using as a substrate maltotriose having a degree of glucose polymerization of 3, the trehabose yield atter enzymatic reaction of a non-reducing seachratef corming enzyme and the present trehalose-releasing enzyme was relatively low, i. e. 4.2%, while in the case of using as a substrate partial starch hydro-lysates having a degree of glucose polymerization of 10-34 1, the trehalose yield was relatively high, i.e. 56.1-80.8%. It was found that the higher the degree of glucose polymerization of material reducing partial starch hydrolysates, the higher the purity of the resultant trehalose. It was also found that the purity of the resultant trehalose and be more increased by allowing glucoamplase to act on a reaction mixture, prepared by the hydrolysis of reducing partial starch

hydrolysates by the two enzymes, to hydrolyze the concomitant non-reducing saccharides, having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher, into trehalose and glucose molecules.

Experiment 6

15

20

25

40

45

Maillard reaction

A solution, containing one % of glycine, 10% of a high-purity trehalose preparation with a purity of 99.5%. d.s.b., obtained by the method in Experiment 4, and 50 mM phosphate bufler (pH 7.0), was kept at 100°C for 90 min, followed by cooling the resultant solution, and determining its absorbance at a wave length of 480 mm in a 1-cm cell. As a control, glucose and mallose were similarly treated as above, and the resultants were subjected to determination of their absorbances at a wave length of 480 mm. The results were as shown in Table 5.

Table 5

Saccharide preparation	Coloration degree (480 nm)
Trehalose (Present invention)	0.006
Glucose (Control)	1.671
Maltose (Control)	0.926

As evident from the results in Table 5. it was revealed that the trehalose preparation according to the present invention only showed a slight coloration induced by the maillard reaction, i.e. the coloration degree was merely about 0.4-0.6% of those of glucose and maltose.

The results showed that the present trehalose is substantially free from the maillard reaction. Thus, the present trehalose is a saccharide which has a less fear of causing deterioration of amino acids when mixed with them.

Experiment 7

Utilization test in vivo

In accordance with the method as reported by Alsuji et al. in "Rinsho-Ejvo", Vol.41, No.2, pp. 200-206 (1972), 30 g of the high-purity trehalose preparation with a purity of 99.5%, d.s.b., obtained by the method in Experiment 4 was dissolved in water into 20 w% % aqueous solution which was then orally administered to 6 healthy maile volunteers, 26-, 27-, 28-, 29-, 30- and 31-year-old. The volunteers were collected their blood at a prescribed time interval, and each collected blood was assayed for its blood sugar- and insulin-levels. As a control, glucose was used. As a result, the trehalose preparation showed the same dynamics as in glucose, i.e. the blood sugar- and insulin-levels showed their maxima at an about 0.5-1 hour after their administrations. It was revealed that the trehalose preparation is readily assimilated, absorbed, metabolized and utilized by the body as an energy source.

Experiment 8

Acute toxicity test

By using mice, the high-purity trehalose preparation with a purity of 99.5%, d.s.b., obtained by the method in Experiment 4 was orally administered to the mice for its acute toxicity test. As a result, it was revealed that the trehalose preparation is a relatively-low toxic substance and no mouse died even when administered with it in an amount of the highest possible dose. Though not so accurate, the LDs $_{00}$ of the trehalose preparation was 50 g/kg or higher.

50 Experiment 9

Production of trehalose-releasing enzyme by Arthrobacter sp. Q36

Similarly as in Experiment 1, a seed culture of Arthrobacter sp. Q36 (FERM BP-4316) was cultured by a fermenter for about 72 hours in place of Arthrobium sp. M-11 (FERM BP-4130). The activities of a non-reducing saccharide-forming onzyme and the present trahalose-releasing onzyme in the resultant culture were respectively about 1.3 units' ml and about 1.9 units'ml. Similarly as in Experiment 1, a cell suspension and a culture supermatant, prepared from the resultant culture, were assayed. The former had an about 0.5 units'ml of non-reducing saccharide-forming activity.

and an about 0.5 units/ml of trehalose-releasing enzyme, while the latter had an about 0.8 units/ml of non-reducing saccharide-forming enzyme and an about 1.3 units/ml of trehalose-releasing enzyme.

Experiment 10

Purification of enzyme

By using an about 18 L of a culture obtained by the method in Experiment 9, the formed non-reducing saccharideforming enzyme was purified similarly as in Experiment 2. The results in each purification step were tabulated in Table 6.

Table 6

Purification step	Enzyme*activity	Specific activity (unit)	Yield (%) (units/mg protein)
Culture	23,700	-	100
Supernatant after cell disruption	22,400	0.15	95
Dialyzed solution after salting out with ammonium sulfate	20,200	0.51	85
Eluate from ion- exchange column	15,100	6.5	64
Eluate from hydrophobic column	8,450	115	36
Eluate from gel filtration column	6,120	217	26

Note: The symbol *** means a non-reducing saccharide-forming enzyme.

25

30

35

40

55

10

15

20

Table 7

Purification step	Enzyme**activity	Specific activity (unit)	Yield (%) (units/mg protein)
Culture	32,500	-	100
Supernatant after cell disruption	30,100	0.19	93
Dialyzed solution after salting out with ammonium sulfate	25,400	0.72	78
Eluate from ion- exchange column	22,700	22.3	70
Eluate from hydrophobic column	15,200	215	47
Eluate from gel filtration column	11,600	497	36

Note: The symbol **** means the present trehalose-releasing enzyme.

Purified enzyme preparations of non-reducing saccharide-forming enzyme and trehalose-releasing enzyme, obtained as the eluates from gel filtration column in Tables 6 and 7, were studied on the purities of the enzymes by using electrophoresis similarly as in Experiment 2. As a result, they were respectively observed as a single protein band, and this meant that they were relatively-high purity enzyme preparations exhibiting an electrophoretically single band.

5 Experiment 11

Property of enzyme

A purified trehalose-releasing enzyme preparation obtained by the method in Experiment 10 was determined its molecular weight on SDS-PAGE to give about 57,000-67,000 dattons. The plot the enzyme preparation was determined in isoelectrophoresis similarly as in Experiment 3 to give a pl of about 3.6-4.6. The influences of temperature and pH on the enzyme activity, as well as the thermal stability and pH stability, were studied similarly as in Experiment 3. The results of the influence of temperature, influence of pH, thermal stability and pH stability were respectively as shown in FIG s 6, 7, 8 and 9.

As evident from these FIG.s, the optimum temperature of the enzyme preparation is about 45° C; the optimum pH, about 6.0-7.5; the thermal stability, up to about 45° C, and the pH stability, about 5.0-10.0.

Experiment 12

Preparation of trehalose from non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher

By using the purified enzyme preparation obtained by the method in Experiment 10, trehalose was experimentally prepared from non-reducing saccharides having a trehalose structure and having a degree of glucose polymerization of 3 or higher according to the methods in Experiment 4. As a result, it was revealed that the enzyme preparation releases trehalose from the non-reducing saccharides similarly as that derived from *Rhizobium* sp. M-11.

Experiment 13

10

15

25

30

35

40

45

50

55

Production and property of trehalose-releasing enzyme from known microorganisms

Among hitherto known microorganisms, those of the species *Brovibactorium helvolum* (ATCC 11822) and *Micrococcus reseus* (ATCC 188), which had been confirmed by the present inventors to produce the present trehalose-releasing enzyme, were respectively cultured by a fermenter at 27°C for 72 hours similarly as in Experiment 1. Eighteen Lof each resultant culture was subjected to a cell disrupting appearative and centrifuged to obtain a supermetant which was then successively salted out with armonium sulfate, dialyzed, and subjected to an ion-ex-braneg column to obtain a partially purified enzyme preparation, followed by studying its properties. The results along with those of *Rhizobium* p. M-11 and *Arthrobactor* p. GS were tabulated in Table 9.

		Table 8		
Microorganism	Enzyme activity of eluate from ion-exchange column (unit)	Optimum temperature (°C)	Optimum pH	Thermal pH Stability (C)
Brevibacterium helvolum (ATCC 11822)	6,070	About 40	About 6.5-6.8	About 6.5-6.8 Up to About 5.5-9.5 about 40
Micrococcus roseus (ATCC 186)	3,010	About 35	About 6.8	Up to About 6.5-7.2 about 30
Rhizobium sp. M-11 (FERM BP-4130)	25,400	About 45	About 6.0-7.5	Up to About 5.0-10.0 about 40
Arthrobacter sp. Q36 (FERM BP-4316)	22,700	About 45	About 6.0-7.5	Up to About 5.0-10.0 about 45

In accordance with the method in Experiment 12, trehalose was experimentally prepared from non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher. As a result, it was revealed that similarly as the trehalose releasing enzyme from *Rhizobium* sp. M-11, every enzyme preparation forms trehalose from the non-reducing saccharides.

Experiment 14

Partial amino acid sequence of trehalose-releasing enzyme

Experiment 14 (1)

15

20

25

30

35

40

Amino acid sequence containing N-terminal

A portion of a purified enzyme preparation derived from Rhizoblums p. M-11, obtained by the method in Experiment 2, and a portion of a purified orzyme preparation derived from Arthobacter sp. O38, obtained by the method in Experiment 10, were dialyzed against distilled water, and about 80µg protein of each resultant preparation was used as a sample for the determination of their amino acid sequences containing their N-terminals. The amino acid sequences were analyzed on "PROTEIN SECUENCER MODEL 473"A, an apparatus of Applied Biosystems, Inc., Foster City, USA, to reveal their 10 amino acid residues from their N-terminals. Partial amino acid sequences containing the N-terminals of the enzyme preparations were as shown in Table 9.

Table 9

	100100		
	Origin	Partial amino acid sequence containing N-terminal	
;	Rhizobium sp. M-11 (FERM BP-4130)	alanine-lysine-proline-valine-glutamine-glycine-alanine-glycine- arginine-phenylalanine	
	Arthrobacter sp. Q36 (FERM BP-4316)	threonine-proline-threonine-tyrosine-proline-arginine-glutamic acid- arginine-alanine-lysine	

As evident from Table 9, it was revealed that the N-terminal of the enzyme preparation from Rhizobium sp. M-11 is alanine which is followed by an amino acid sequence of lysine-proline-valine-glutamine-glycine-arginine-phenylalanine. In the case of the enzyme preparation from Arthrobactor sp. Q36, the N-terminal is throonine which is followed by an amino acid sequence of proline-throonine-tyrosine-proline-arginine-glutamic acid-arginine-lanine-lysine.

Experiment 14 (2)

Internal partial amino acid sequence

A portion of a purified enzyme preparation derived from Rhizobium sp. M-11, obtained by the method in Experiment 2, and a portion of a purified enzyme preparation derived from Arthrobacter sp. Q36, obtained by the method in Experiment 10, were dialyzed against 10 mM Tris-HCl buffer (pH 9.0), and the resultants were respectively diluted with a fresh preparation of the same buffer to give a concentration of about one mg/ml. To one ml aliquot of the resultant each solution was added 10µg "LYSYL ENDOPEPTIDASE", a product of Wako Pure Chemical Industries, Ltd., Tokyo, Japan, and allowed to react at 30°C for 22 hours to form peptides which were then separated on reverse phase highperformance liquid chromatography (reverse phase HPLC). The apparatus and conditions used to separate the pertides of the enzyme preparation from Rhizobium sp. M-11 in the reverse phase HPLC were "CAPCELL PAK C18 column", a diameter of 4.6 mm and a length of 250 mm, a product of Shiseido Co., Ltd., Tokyo, Japan; a flow rate, 0.6 ml/min, a temperature, an ambient temperature; an elution time, 60 min; and a gradient, a liner gradient of a solution containing 0.1 v/v % trifluoro acetate and acetonitrile ranging from 16-48 v/v %. The apparatus and conditions used to separate the peptides of the enzyme preparation from Arthrobacter sp. Q36 in the reverse phase HPLC were "µ-BONDAPAK C18 column* having a diameter of 2.1 mm and a length of 150 mm, a product of Waters Chromatography Div., MILLIPORE Corp., Milford, USA; a flow rate, 0.9 ml/min; a temperature, an ambient temperature; an elution time, 60 min; and a gradient, a liner gradient of a solution containing 0.1 v/v % trifluoro acetate and acetonitrile ranging from 30-55 v/v %. The peptides eluted from the columns were detected by monitoring an absorbency at a wavelength of 210 nm. From the enzyme preparation of Rhizobium sp. M-11 three peptides named as RT41, RT46 and RT54 having respective retention times of about 41, 46 and 54 were separated from other concomitant peptides, and from the

enzyme preparation of Arthrobactersp. O36 three pepticles named as AT7. AT30 and AT46 having respective retention times of about 7, 30 and 48 were separated from other concomitant peptides, followed by drying the separated peptides in vacuo and dissolving the resultants in 200 µl solutions having different concentrations of 0.1-50 vv %, acetonitrile Each peptide specimen thus obtained was analyzed on a protein sequencer for its amino acid sequence up to 10 amino acid residues. The analyzed internal partial amino acid sequences were as shown in Table 10.

		Table 10
Origin	Peptide	Internal partial amino acid sequence
Rhizobium sp. M-11 (FERM BP-4130)	RT41	histidine-glycine-glutamic acid-glycine-asparagine- threonine-tryptophane-glycine-aspartic acid-serine
	RT46	aspartic acid-glutamic acid-argınine-alanine-valine-histidine- isoleucine-leucine-glutamic acid-glutamic acid
	RT54	leucine-aspartic acid-tryptophane-alanine-glutamic acid- alanine-serine-alanine-glycine-aspartic acid
Arthrobacter sp. Q36 (FERM BP-4316)	AT30	glutamine-glycine-glutamic acid-glycine-asparagine- threonine-tryptophane-glycine-aspartic acid-serine
	AT48	aspartic acid-glutamic acid-arginine-alanine-valine-histidine- isoleucine-leucine-glutamic acid-aspartic acid
	AT7	leucine-aspartic acid-tryptophane-alanine-glutamic acid- alanine-alanine-glutamic acid-glycine-aspartic acid

As ovident from Table 10, 9 of 10 amino acid residues in the internal partial amino acid sequence of peptide RT41 of the enzyme derived drom Rhzobulum sp. M-11 connoided with those of peptide AT30 of the enzyme derived from Athrobacter sp. C35, while those of peptide RT46 coincided with those of peptide AT34 with respect to their 9 of 10 amino acid residues. In the case of peptide RT46 coincided with those of peptide AT48 with respect to their 8 of 10 amino acid residues. Accordingly, it is judged that the enzyme derived from microorganisms of the genus Athrobacter have a relatively-high homology with respect to their internal derived from microorganisms of the genus Athrobacter have a relatively-high homology with respect to their internal partial amino acid sequences which can be expressed by leucine-asparagine-tryptophane-alanine-glutamic acid-alanine-X,-X₂-glycine-aspartia caid (where "X," means "serine" or "alanine", and "X₂" means "slainine" or "glutamic acid-glutamic acid-grainine-alanine-valine/alanine-sloudine-loucine-glutamic acid-X₂ (where "X₃" means "glutamic acid-grainine-alanine-valine-alanine-alanine-sloudine-glutamic acid-X₃ (where "X₃" means "glutamic acid-grainine-alanine-valine-alanine-alanine-sloudine-glutamic acid-X₃ (where "X₃" means "glutamic acid-grainine-alanine-valine-alanine-alanine-sloudine-glutamic acid-X₃ (where "X," means "sloudine-asparagine-throonine-tryptophane-gly-cine-asparagine-serine (where "X," means "sloudine" or "glutamic acid-grainine-alanine-sloudine-glutamic acid-x₃ (where "X," means "sloudine" or "glutamic acid-grainine-alanine-sloudine-glutamic acid-ycine-asparagine-throonine-tryptophane-gly-cine-asparagine-serine (where "X," means "sloudine" or "glutamic acid-grainine-alanine-sloudine-gly-cine-asparagine-throonine-tryptophane-gly-cine-asparagine-serine (where "X," means "sloudine" or "glutamic acid-grainine-alanine-glutamic acid-grainine-alanine-glutamic acid-grainine-alanine-glutamic acid-grainine-alanine-glutamic acid-grainine-alanine-glutamic

The following Examples A illustrate the preparation of the present trehalose-releasing enzyme, trehalose prepared therewith, and saccharide compositions containing the trehalose; and Examples B illustrate compositions containing one or more of these trehalose and saccharide compositions:

Example A-1

10

15

20

25

A seed culture of Intizoblumsp, M-11 (FERM BP-4130) was inoculated in a nutrient culture medium and incubated by a fermenter for about 80 hours in accordance with the method in Experiment 1. After completion of the incubation, the resultant culture was filtered to remove cells with an SF-membrane to obtain an about 18 L filtrate, followed by concentrating the filtrate with a UF-membrane into an about one L enzyme solution containing 17.2 unitsful of a non-reducing associaride-forming onzyme and 20.8 unitsful of the respect trefalose-releasing conzerved.

To 15% suspension of potate starch, d.s.b., was added calcium carbonate to give a final concentration of 0.1% d.s. h, and the resultant solution was adjusted to pH 6.0, mixed with 0.2% by weight of TERMAMYL 6.0.1, ance-amylase specimen commercialized by Novo Industri A/S, Copenhagen, Denmark, and subjected to an enzymatic reaction at 95°C for 15 min. The reaction mixture was autoclaved at a pressure of 2 kg/cm² for 30 min, cooled to 45°C, admissible with 1,000 unities graterin, d.s.b., of "PULLULANSE (EG. 2.2.1 H)", a enzyma specimen commercialized by Hayashibara Biochemical Laboratories, inc., Okayama, Japan, and 0.2 ml/g starch, d.s.b., of the above enzyme solution, and subjected to an enzymatic reaction for 48 hours. The reaction mixture thus obtained was kept at 95°C for 10 min, cooled and filtered, and the resultant filtrate was in usual manner decolored with an activated charcoal, desaited and purified with ion-exchangers in H- and OH-form, followed by concentrating the resultant solution to obtain a 50% syrup in a yield of about 52%, d.s.b.

The syrup, which contains 70.2% trehalose, 2.4% glucosyltrehalose, 3.3% maltosyltrehalose, 0.7% glucose, 10.1%

mallose, 12.9% mallotriose and 0.4% of mallotetraose and higher oligosaccharides, d.s.b., has a moderate and highquality sweetness and a relatively-low reducing power, as well as an appropriate viscosity and moisture-retaining ability. Because of these it can be arbitrarily used in lood products, cosmetics and pharmaceuticals as a sweetener, tasteimproving agent, quality-improving agent, stabilizer, diluent, filter and excipient.

Example A-2

A saccharide solution as a feed solution, obtained by the method in Example A-1, was fractionated by using a column packed with XT-1016 (Ma-1-orm, polymerization degree of 4%); an alkaline-metal strong-aci dation exchange resin commercialized by Tokyo Organic Chemical Industries Ltd, Tokyo, Japan. The procedure was as follows: The resin was packed in 4-jacketed stainless steel columns having an inner diameter of 5.4 cm, and the columns were cascaded in series to give a total gel-bed depth of 20 m. The columns were heated to give the inner column temperature of 55°C and fed with 5 w/ % of the saccharide solution and to remove concomitant saccharides such as mallose and maltotriose, followed by recovering trehalose-rich fractions. The fractions thus obtained were pooled, purified, concentrated, frield in vacure and pulverized to obtain a high trehalose content prowder in a yield of about 55°C, he

The content of trehalose in the product is about 97%, d.s.b., and the product has a mild and high-quality sweetness, and, because of these it is arbitrarily used in food products, cosmetics and pharmaceuticals as a sweetener, taste-improving agent, quality-improving agent, stabilizer, excipient and filler.

Example A-3

15

20

25

40

45

50

55

A high trehalose content fraction obtained by the method in Example A-2 was in usual manner decolored with an activated charcoal, desalled with an ion-exchanger, and concentrated into an about 70% solution which was then placed in a crystallizer, admixed with about 2% hydrous crystalline trehalose as a seed crystal, and gradually cooled to obtain a massecutie with a crystallinity of about 45%. The massecutie was sprayed from a nozzle equipped at the top of a drying lower at a high pressure of 150 kg/cm², in the spraying step, the massecutie was simultaneously ventilated with 35°C hot air being sent from the top of the drying lower, and the resultant crystalline powder was collected on a metal wire netting conveyer provided on the basement of the drying lower, and gradually moved out of the drying tower, and gradually noved out of the drying tower, and gradually noved out of the drying tower while a stream of 45°C air was passing upwards through the metal wire netting. The resultant crystalline powder was injected in an ageing tower and aged for 10 hours to complete the crystallization and drying, followed by recovering a powdery hydrous crystalline trehalose in a yield of about 90% against the material high trehalose content fraction, ds.b.

The product is substantially non-hygroscopic and handles easily, and these render it arbitrarily useful in food products, cosmetics and pharmaceuticals as a sweetener, taste-improving agent, quality-improving agent, stabilizer, excipient and filler.

Example A-4

A high trehalose content fraction obtained by the method in Example A-2 was purified smillarly as in Example A-3 and the resultant was placed in an evaporation, and boiled up in vacuo to obtain a syrup with a molisture content of about 3.0%. The resultant syrup was placed in a crystallizer, admixed with one % anhydrous crystalline trehalose against the dry weight of the syrup, and crystallized at 120°C for 5 min under stirring conditions, and the resultant mixture was placed in a alluminum plain container and aged at 100°C for 6 hours to obtain a blown of the syrup.

The resultant block was pulvarized by a cutter and dried by a fluidized-bed drying to obtain a powdery anhydrous crystalline trehalose with a moisture content of about 0.3% in a yield of about 85% against the material high trehalose content fraction, d s b. The product can be arbitrarily used as a desiccant in food products, cosmetics, pharmaceuticals, and their materials and intermediates, and also can be used as a white powdery sweetener in a variety of compositions such as food oroducts, cosmetics and observaceuticals.

Example A-5

In accordance with the method in Example A-1, a seed culture of a mutant of *Rhizobium* sp. M-11 (FEFM BP-4130) was inoculated in a nutrient culture medium and incubated by a termenter for about 70 hours. After completion of the incubation, the resultant cells were membrane filtered with a SF-membrane to recover an about 100. filtrate which was then concentrated with a UF-membrane to obtain an about 51 cnzyme solution containing about 410 units/mid of anon-reducing ascharide-forming enzyme, and about 440 units/mid for therelases-releasing enzyme.

Six % suspension of potato starch was gelatinized by heating, adjusted to pH 4.5, heated to 50°C, admixed with

500 units/g starch, d.s.b., of an isoamylase specimen commercialized by Hayashibara Biochemical Laboratories, Inc., Okayama Japan, and subjected to an enzymatic reaction for 20 hours. The reaction mixture was adjusted to pH 6.5. autoclaved at 120°C for 10 min, cooled to 95°C, admixed with 0.1% per g starch, d.s.b., of "TERMAMYL 60L", an αamylase specimen commercialized by Novo Industri A/S Copenhagen Denmark, and subjected to an enzymatic reaction for 15 min. The reaction mixture was autoclaved at 130°C for 30 min, cooled to 45°C, admixed with 0.01 ml per q starch, d.s.b., of the above-mentioned enzyme solution, and subjected to an enzymatic reaction for 64 hours. The resultant reaction mixture was kept at 95°C for 10 min, cooled to 50°C, adjusted to pH 5.0, admixed with 10 units/g starch, d.s.b., of "GLUCOZYME", a glucoamylase specimen commercialized by Nagase Biochemicals, Ltd., Kyoto, Japan, subjected to an enzymatic reaction for 40 hours, and heated to inactivate the remaining enzyme. The solution thus obtained was in usual manner decolored with an activated charcoal, desalted with an ion-exchanger and concentrated into an about 60% solution containing 80.5% trehalose, d.s.b. The solution was concentrated into an about 84% solution which was then placed in a crystallizer and admixed with an about 2% hydrous crystalline trehalose as a seed crystal against the dry weight of the solution to crystallize trehalose under stirring conditions. The resultant was placed in a plain plastic-vessel and allowed to stand at an ambient temperature for 3 days to form a block which was then pulverized by a cutter to obtain a powdery hydrous crystalline trehalose in a yield of about 90% against the material starch, d.s.b.

The product is substantially non-hygroscopic and handles easily, and these render it arbitrarily useful in a variety of compositions such as food products, cosmetics and pharmaceuticals as a sweetener, taste-improving agent, quality-improving agent, stabilizer, excipient and filler.

Example A-6

15

20

25

30

45

50

55

A seed culture of a microorganism of Arthrobacter sp. Q36 (FERM BP-4316) was inoculated in a nutrient culture medium and cultured with a farmenter for about 72 hours in accordance with the method in Experiment 9. The resultant culture was centrifuged at 10,000 pm for 30 min to remove cells, and the resultant supernatant was concentrated with a IP-membrane to obtain one L of an enzyme solution containing 16.3 units/ml of a non-reducing saccharide-forming enzyme and 25.1 units/ml of the present trehabes-releasing enzyme.

One part by weight of potato starch was mixed with 6 parts by weight of water and 0.01 part by weight of "NEO-SPITASE*, an α-amylase specimen commercialized by Nagase Biochemicals Ltd., Kyoto, Japan, and the resultant mixture was mixed by stirring, adjusted to pH 6.2 and heated to 85-90°C to gelatinize and liquefy the starch. The resultant liquefied starch was autoclaved at 120°C for 10 min to inactivate the remaining enzyme, cooled to 45°C. admixed with 500 units/g starch of an isoamylase specimen commercialized by Hayashibara Biochemical Laboratories, Inc., Okayama, Japan, and 0.2 ml/g starch of the above enzyme solution, and subjected to an enzymatic reaction for 48 hours. After completion of the reaction, the reaction mixture was heated at 95°C for 10 min to inactivate the remaining enzyme, adjusted to 50°C and pH 5.0, admixed with 10 units/g starch "GLUCOZYME", a glucoamylase specimen commercialized by Nagase Biochemicals Ltd., Kyoto, Japan, subjected to an enzymatic reaction for 40 hours, and heated to inactivate the remaining enzyme. The reaction mixture thus obtained was in usual manner decolored with an activated charcoal, desalted with an ion-exchanger, and concentrated into an about 60% solution containing 78.3% trehalose, d.s.b. In accordance with the method in Example A-2 except for using as the ion-exchanger "CG 6000 (Na*form)", an alkaline-metal strong-acid cation-exchange resin commercialized by Japan Organo Co, Ltd., Tokyo, Japan, the concentrated solution was subjected to an ion-exchange column chromatography to obtain a high trehalose content fraction containing about 95% trehalose, d.s.b. The fraction was concentrated to give a concentration of 75%, placed in a crystallizer, crystallized by the addition of about 2% hydrous crystalline trehalose as a seed crystal under stirring conditions, transferred to a plain plastic-container, allowed to stand and aged at an ambient temperature for 3 days to obtain a block, followed by pulverizing it with a cutter to obtain a powdery hydrous crystalline trehalose in a yield of about 70% against the material starch, d.s.b.

The product, which is substantially free from hygroscopicity and handles easily, can be arbitrarily used as a sweetener, taste-improving agent, quality-improving agent, excipient, filler and diluent in a variety of compositions such as food products, cosmetics and pharmaceuticals.

Example A-7

In accordance with the method in Experiment 13, a seed culture of a microorganism of the species *Brevibacterium* helvolum (ATCC 11822) was incoulated in a nutrient culture medium, and cultured by a fermenter for about 72 hours, and the resultant culture was treated with a cell disrupting apparatus. The resultant mixture was contributed 10,000rpm for 30 min to remove residues, and the resultant supernatant was concentrated with a UF-membrane, followed by the recovery of an about 720 ml solution containing about 8 units/ml of a non-saccharide-forming enzyme and about 120 units/ml of a technique-releasing enzyme.

A 33% tapicae starch suspension was admixed with calcium carbonate to give a final concentration of 0.1%, adjusted to pl R.0. admixed with 0.3% "TERMAN"L 6.0L", an c-ampliase specimen commercialized by Novo industria A/S, Copenhagen, Denmark, and subjected to an enzymatic reaction at 95°C for 20 min. The resultant reaction mixture was autoclaved under a pressure of 2 kg/cm² for 30 min. cooled to 40°C, admixed with 200 units/g starch of an iso-amylase specimen commercialized by Hayashibara Biochemical Laboratories, inc., Okayama, Japan, and 0.2 mig starch of the above enzyme solution, and subjected to an enzymatic reaction for 48 hours. The reaction mixture thus obtained was kept at 95°C for 10 min, cooled and filtered to obtain a filtrated with was then in usual manner decode with an activated charcoal, desalted with ion-exchangers in H- and OH-form, and concentrated to obtain a 60% syrup in a viel of a hour 190%, d. 5

The product, which contains 60.1% trehalose, 1.4%, glucosyltrehalose, 1.5% mallosyltrehalose, 1.0% glucose, 6.5% maltose, 8.3% maltotriose, 2.1.2% maltotetraces and higher maltooligosaccharides, has a mild and high-quaitly sweatness as well as a relatively-low reducing power and an adequate moisture-retaining ability. Because of these, it can be used in a variety of compositions such as food products, cosmotics and pharmaceuticals as a sweetener, taste-improving agent, sabilityer, oxiciplent, filter and dilluent.

Example A-8

15

35

40

45

50

A high trehalose content solution containing about 95% trehalose, obtained by the method in Example A-6, was in usual manner decolored and desalted. The resultant solution was concentrated into an about 75% solution which was then transferred to a crystallizer, admixed with about 2% hydrous crystalline trehalose as a seed crystal, heated to 50°C, gradually cooled to 25°C while stirring, and subjected to a separation using a basket-type centrifuge. The resultant crystal was washed by spraying it with a small amount of water to obtain a high-purity hydrous crystalline trehalose with a purity of 99% or higher in a yield of about 50%.

25 Example B-1

Sweetener

To one part by weight of a powdery hydrous crystalline trehalose, obtained by the method in Example A-5, was homogeneously added 0.01 part by weight of "αS SWEET", an α-glycosyl stevioside product commercialized by Toyo Sugar Refining Co., Ltd., Tokyo, Japan, and 0.01 part by weight of "ASPAFTAME", a product of L-aspartyl-phenylalanine methylester commercialized by Ajinomoto Co., Ltd., Tokyo, Japan, and the resultant mixture was fed to a granulator to obtain a granular sweetener. The product has a satisfactory sweetness and an about 2.5-fold higher sweetening power of sucrose, and the caloric value is lowered to about 2/5 of that of sucrose.

Since the product has a satisfactory stability and does not decompose other sweeteners to be mixed, it can be suitably used as a low-caloric sweetener for low-caloric food products for fat persons and diabetics who are restricted to a reduced calorie intake.

The product substantially does not form acids and insoluble glucans when dental carries-inducing microorganisms act on it, and this renders it useful for sweetening food products to prevent dental carries.

Example B-2

Hard candy

One hundred parts by weight of 55% sucrose solution was mixed while heating with 30 parts by weight of a trehalose syrup, obtained by the method in Example A-7, and the resultant solution was concentrated in vacuo until the moister content lowered to below 2%. The concentrated solution was admixed with one part by weight of clinc acid and adequate amounts of a lemon flavor and a coloring agent, and the resultant mixture was in usual manner formed into the desired product.

The product is a high-quality hard candy having a satisfactory taste and biting property, as well as having no fear of causing crystallization of sucrose.

Example B-3

55 Chewing gum

Three parts by weight of a gum base was melted by heating until it softened, and the resultant was mixed with 4 parts by weight of sucrose and 3 parts by weight of a hydrous crystalline trehalose powder obtained by the method in

Example A-3, and further mixed with adequate amounts of a flavor and a coloring agent. The resultant mixture was in usual manner kneaded by a roll, formed and packed to obtain the desired product

The product is a chewing gum having a satisfactory texture and taste.

Example B-4

Sweetened condensed milk

Three parts by weight of a trehalose syrup obtained by the method in Example A-1 and one part by weight of sucrose were dissolved in 100 parts by weight of fresh milk, and the resultant solution was sterilized by heating with a plate heater, and condensed to give a concentration of 70%, followed by aseptically canning the resultant concentrate into the desired product.

The product has a mild sweetness and a satisfactory taste, and these render it arbitrarily useful as a seasoning for baby foods, foods for infants, fruit, coffee, cocoa and tea.

Example B-5

15

20

25

Lactic acid beverage

One hundred and seventy-five parts by weight of defatted milk. 190 parts by weight of a trehalces syrup obtained by the method in Example A.1, and 50 parts by weight of a high tactosucrose content powder as disclosed in Japanese Patent Laid Open No.281.795/92 were dissolved in 1.150 parts by weight of water. The resultant solution was sterilized at 65°C for 30 min, cooled to 40°C and inoculated with 30 parts by weight of lactic acid bacterium as a starter, followed by the incubation at 37°C for 8 hours to obtain the desired product.

The product is a lactic acid beverage with a satisfactory taste and flavor. Since the product contains oligosaccharides, it stably retains lactic acid bacteria and promotes the growth of bifid bacteria.

Example B-6

30 Powdered juice

Thirty-three parts by weight of a powdered orange juice prepared by spray drying was mixed to homogeneity with 50 parts by weight of a high trehalese content prowder obtained by the method in Example A.2. In parts by weight of sucrose, 0.85 parts by weight of anhydrous citric acid, 0.1 part by weight of malic acid, 0.1 part by weight of anhydrous citric acid, 0.1 part by weight of malic acid, 0.1 part by weight of anhydrous citric acid, 0.1 part by weight of malic acid, 0.1 part by weight of a powdered acid. 0.1 part by weight of weight of publish, and an adequate amount of a powdered acid. Or part by weight of malic acid, 0.1 part by weight of a part by weight of publish, and an adequate amount of a powdered acid. Part of parts of

The product containing 30% orange juice, d.s.b., retained its high quality for a relatively-long period of time without giving an unsatisfactory taste and smell.

Example B-7

Custard cream

One hundred parts by weight of com starch, 100 parts by weight of a trehalcse syrup obtained by the method in Example A.7, 80 parts by weight of maltose, 20 parts by weight of sucrose, and one part by weight of salt were mixed to homogeneity. The resultant mixture was admixed with 280 parts by weight of egg, and gradually added with 1,000 parts by weight of a boiling milk. The mixture thus obtained was continued stirring under heating orangitions, and the heating was stopped when the com starch in the mixture was completely gelatinized to give the whole contents semitransparent, followed by cooling the mixture and adding thereto an adequate amount of a vanilla flavor. The resultant mixture was weighted, injected and packed to obtain the desired product.

The product has a smooth surface and gloss, as well as a mild taste and sweetness.

Example 8

Uiro-no-moto (premix of starch paste)

Ninety parts by weight of rice powder, 20 parts by weight of com starch, 40 parts by weight of sucrose, 80 parts by weight of a powdery hydrous crystalline trehalose obtained by the method in Example A-3, and 4 parts by weight of pullulan were mixed to homogeneity to obtain a *ution-on-moto*. The product was kneaded with adequate amounts of matchs (green tea) and water, and the mixture was placed in a container and steamed up for 60 min to obtain a *matcha-utio*.

The product has a satisfactory gloss and biting property, as well as a satisfactory flavor and taste, and has a relatively-long shelf-life because the retrogradation of starch is well inhibited.

Example B-9

10

25

15 An (beans paste)

Ten parts by weight of adzuki beans as a material was in usual manner mixed with water and boiled, followed by removing the astringency, harshness of the beans, and water-soluble impurities to obtain about 21 parts by weight of "adzuki-subu-an". To the resultant were added 14 parts by weight of sucrose, 5 parts by weight of a trehalose syrup obtained by the method in Example A-1, and 4 parts by weight of water, and the resultant mixture was boiled, mixed with a small amount of salad oil, and carefully kneaded up so as not to paste the beans. Thus, the desired product was obtained in a visid of about 35 kg.

The product free from discoloration induced by boiling has a satisfactory taste and flavor, and these render it useful as a material an for bean-jam buns, buns with bean-jam filling, dumplings, bean-jam-filled wafers, sherbets and be creams

Example B-10

Bread

One hundred parts by weight of wheat powder, 2 parts by weight of yeast, 5 parts by weight of sugar, one part by weight of a powdery hydrous crystalline terhalose obtained by the method in Example A-3, 0.1 part by weight of inorganic yeast food were kneaded with water in usual manner, femented at 26°C for 2 hours, aged for 30 min and baked up.

The product is a high-quality bread having a satisfactory hue and rising, as well as a satisfactory elasticity and mild sweetness.

Example B-11

Ham

45

To one thousand parts by weight of sliced ham meat were added and ground to homogenelly 15 parts by weight of salt and 3 parts by weight of potassium nitrate, and the ham meat slices were piled up and allowed to stand overnight in a cold-storage room. Thereafter, the resultant slices were first scaked in a salt solution consisting of 500 parts by weight of water, 100 parts by weight of salt. 3 parts by weight potassium nitrate, 40 parts by weight of a powder containing non-reducing saccharidates prepared by the method in Example A-6, and an adequate amount of a peperint for 7 days in a cold-storage room, then washed with cold water in usual manner, tied up, smoked, cooked cooled and packed to obtain the desired product.

The product is a high-quality ham having a satisfactory hue, taste and flavor.

50 Example B-12

Powdery peptide

One part by weight of "HINUTE S*, a peptide solution containing 40% edible soy beans commercialized by Fuji Oil Co., Ltd., Tokyo, Japan, was mixed with 2 parts by weight of a powder containing a powdery hydrous crystalline trehalose prepared by the method in Example A-6, and the resultant mixture was placed in a plastic vessel, dried in vacuo at 50°C, and pulverized to obtain a powdery postide.

The product having a satisfactory taste and flavor can be arbitrary used as a material for confectioneries such as

premixes, sherbets and ice creams, as well as baby foods and nutritions for therapy in the form of an oral or an intubation feeding.

Example B-13

Powdered miso

To one part by weight of akamiso (a kind of miso) was added 3 parts by weight of a powdery anhydrous crystalline trehalose obtained by the method in Example A-4, and the mixture was poured into a metal plate having hemisphere wells on its surface and allowed to stand at an ambient temperature overnight to obtain miso solide, about 4 g weight each, which were then subjected to a pulverizer to obtain the desired product.

The product can be arbitrarily used as a seasoning for instant noodles and soups, as well as a miso confectionery.

Example B-14

15

25

40

45

55

Powdery egg volk

Egg yolks prepared from fresh eggs were sterilized at 60-64°C by a plate heater, and one part by weight of the resultant liquid was mixed with 4 parts by weight of a powdery anhydrous crystalline trehalose prepared by the method in Example A-4 with respect to one part by weight of the liquid. The resultant mixture was transferred to a vessel, allowed to stand overnight to form a block while the anhydrous crystalline trehalose was allowing to convert into hydrous crystalline trehalose. The block thus obtained was pulverized by a cutter to obtain a powdery egg yolk.

The product can be arbitrarily used as a material for confectioneries for premixes, sherbets, ice creams and emulsifiers, as well as bably foods and nutritions for therapy in the form of an oral or an intubation feeding. The product can be also used as a skir refiner and hair restorer.

Example B-15

Cosmetic cream

Two parts by weight of polyoxyelrylene glycol monostearate, 5 parts by weight of glyceryl monostearate, selfemulaliying, 2 parts by weight of a high trehalese content powder obtained by the method in Example A.2, one part by weight of or-glycosyl rutin, one part by weight of liquid petrolatum, 10 parts by weight of glyceryl tri-2-ethylhexancate, and an adequate amount of an antiseptic were in usual manner dissolved by heating. The resultant solution was admixed with 2 parts by weight of 1-gast by weight of 1-gast by weight of 10 parts by weig

The product having a relatively-high stability can be arbitrarily useful as a high-quality sunscreen, skin-refining agent and skin-whitening agent.

Example B-16

Powdery ginseng extract

A half part by weight of ginseng extract was mixed with 1.5 parts by weight of a powdery anhydrous crystalline trehalose prepared by the method in Example A-4, and the resultant mixture was transferred to a plain container, allowed to stand for 2 days to convert anhydrous crystalline trehalose into hydrous crystalline trehalose to form a block. The resultant block was pulverized by a cutter and classified to obtain a powdery ginseng extract.

The product and adequate amounts of powdery vitamins B1 and B2 were subjected to a granulator to obtain a powdery ginseng extract containing vitamins.

The product thus obtained can be arbitrarily used as a tonic, fatigue-relieving agent and vitality-imparting agent. The product can be also used as a hair restorer.

Example B-17

Solid pharmaceutical

A natural human interferon-α preparation, commercialized by Cosmo Bio, Tokyo, Japan, was in usual manner fed

to a column of an immobilized anti-human interferon-a entibody to adsorb the interferon-a, and a buffer containing calf serum albumin as a stabilizer was fed to the column, followed by removing an excessive amount of the albumin. Thereafter, the interferon-a was eluted from the column with a physiological saline containing 5% of a high trehalose content powder, prepared by the method in Example A-2, while the pH of the physiological saline was varying. The resultant foliate was membrane filtered, and the filtrate was dehydrated by the addition of about 20-fold volume of "FINETOSE®", an anhydrous crystalline maltose powder commercialized by Hayashibara Shoji, Inc., Okayama, Japan, followed by pulverizing the resultant todawing the resultant todawing membrane to obtain tablets containing about 150 units of the natural human interferon-a per one tablet, about 200 mg weight.

The product can be orally administered as a sublingual tablet to patients at a dose of 1-10 lablets/edult/day, and arbitrarily used to treat viral diseases, allergys, rheumatisms, diabetes and malignant tumors. More particularly, the product can be suitably used as a therapeutic agent for AIDS and hepatitis, the number of patients suffering from these diseases has been remarkably increased. The trehalose and maltose incorporated in the product act as a stabilizer for the natural human interferon-or, so that the activity is well retained for a relatively-long period of time even at an ambient temperature.

Example B-18

15

20

25

30

35

40

45

50

55

Sugar coated tablet

A crude tablet as a core, 150 mg weight, was coated with a solution consisting of 40 parts by weight of a powdery hydrous crystalline trehalose obtained by the method in Example A-3, 2 parts by weight of pullulan having an average molecular weight of 200,000, 30 parts by weight of water, 25 parts by weight of falc, and 3 parts by weight of Itanium oxide until the total weight reached to about 230 mg, and the resultant was further coated with a solution consisting of 5 parts by weight of affects preparation of the same powdery hydrous crystalline trehalose, one part by weight of pullulan, and 34 parts by weight of water, and glossed with a liquid wax to obtain a sugar coated tablet having a satisfactory gloss and appearance.

The product has a relatively-high shock tolerance and retains its high quality for a relatively-long period of time.

Example B-19

Dentifrice

A dentifrice was prepared in usual manner by mixing the following ingredients:

Calcium monohydrogenphosphate	45.0%
Pullulan	2.95%
Sodium lauryl sulfate	1.5%
Glycerine	20.0%
Polyoxyethylene sorbitan laurate	0.5%
Antiseptic	0.05%
Powdery hydrous crystalline	12.0%
trehalose prepared by the	
method in Example A-3	
Maltitol	5.0%
Water	13.0%

The product is satisfactorily used as a dentifrice for infants because it has an adequate sweetness.

Example B-20

Solid preparation for intubation feeding

A composition consisting of the following compositions was prepared. Five hundred parts by weight of a powdery hydrous crystalline trehalose prepared by the method in Example A6, 270 parts by weight of powdered egg yolk, 209 parts by weight of defatted milk, 4.4 parts by weight of defatted milk, 4.4 parts by weight of sodium chloride, 1.8 parts by weight of sodium ascorbate, 0.6

parts by weight of vitamin E acetate, and 0.04 parts by weight of nicotinamide. Twenty-five g aliquots of the composition were injected into moisture-proof laminated small bags and heat sealed to obtain the desired product.

One bag of the product is dissolved in about 150-300 ml of water into a fluid food, and orally or parenterally administered to nasal cavity, stomach or intestine by intubation feeding to supplement energy to living bodies.

Example B-21

Hyperalimentation

A high-purity hydrous crystalline trehalose, prepared by the method in Example A-8, was dissolved in water into an about 10 wiv % aqueous trehalose solution which was then in usual manner membrane filtered to remove pyrogen, assotically injudiced into a plastic bottle, and sealed to obtain the desired product.

The product, which is a satisfactorily stable hyperallimentation substantially free of change on standing, is suitable for intravenous- and intrapentoneal-administrations. A 10 w/v % solution of the product is isotonic to blood, and this means it can supplement energy to living bodies at 2-fold higher concentration than in the case of cluoses.

Example B-22

15

20

25

30

35

40

45

Hyperalimentation

A high-purity hydrous crystalline trehalose, prepared by the method in Example A-8, and an amino acid composition consisting of the following components were dissolved by stirring in water to give respective concentrations of 5 w/v % and 30 w/v %, and, similarly as in Example B-10 the resultant solution was purified to obtain a pyrogen-free solution, followed by injecting it into a plastic bottle and sealed to obtain the desired product.

Components of amino acid co	mposition
Component	mg/100 ml
L-Isoleucine	180
L-Leucine	410
L-Lysine monohydrochloride	620
L-Methionine	240
L-Phenyl alanine	290
L-Threonine	180
L-Tryptophane	60
L-Valine	200
L-Arginine hydrochloride	270
L-Histidine monohydrochloride	130
Glycine	340

Although the product is a multiple hyperalimentation containing trehalose and amino acids, it is satisfactorily stable without substantial change on standing and can be suitably administered intravenously and intraperitoneally to living bodies. The product can be arbitrarily used to supplement energy as well as amino acids to living bodies.

50 Example B-23

Ointment for treating trauma

Two hundred parts by weight of a high trehalose content powder, prepared by the method in Example A-2, and 300 parts by weight of mallose were admixed with 50 parts by weight of methanol solution containing 3 parts by weight of iodine, and the resultant solution was mixed with 200 parts by weight of a 10 wb % aqueous pullulan solution to obtain the desired product having a satisfactory extensibility and adhesiveness.

The iodine contained in the product exerts a bactericidal activity, and the trehalose in the product acts as an energy-

supplementing agent on viable cells, and because of these the product shortens a healing period and satisfactorily heals a wound surface.

As is evident from above, the present novel trehalose-releasing enzyme releases trehalose from non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher. and forms trehalose in a relatively-high yield when allowed to act on reducing partial starch hydrolysates together with a non-reducing saccharide-forming enzyme. The trehalose thus obtained can be readily separated and purified, and the resultant purified trehalose and saccharide compositions containing it has a satisfactory stability as well as a relatively-high quality and mild sweetness. It is readily assimilated, absorbed and utilized by living bodies when orally administered intact or parenterally administered in the form of a transfusion agent. Trehalose per se and saccharide compositions containing the same can be arbitrarily used as a sweetener, taste-improving agent, quality-improving agent, stabilizer, excipient and filler in a variety of compositions such as food products, cosmetics and pharmaceuticals.

Thus, the present invention provides a novel technique to prepare trehalose and saccharide compositions containing the same in an industrial-scale and a relatively-low cost from partial starch hydrolysates prepared from starch. a cheap and abundant natural source. Therefore, the present invention gives an unfathomable great influence on the fields such as starch-, enzyme- and biochemical-sciences, and other industrial fields, especially, food-, cosmetic- and pharmaceutical-industries, as well as forestry, fisheries, and agricultural-, livestock- and chemical-industries. Thus, the influence of the present invention on the fields is unfathomably great.

While there has been described what is at present considered to be the preferred embodiments of the invention, it will be understood the various modifications may be made therein, and it is intended to cover in the appended claims all such modifications as fall within the true spirits and scope of the invention.

Claims

20

30

35

40

45

50

- 25 1. A trehalose-releasing enzyme which specifically hydrolyzes the linkage between a trehalose moiety and the remaining glycosyl moiety in a non-reducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher.
 - 2. An enzyme according to claim 1, wherein said glycosyl moiety consists of one or more glucose residues.
 - 3. An enzyme according to claim 1 or claim 2, which has the following physicochemical properties:
 - (1) Action
 - Specifically hydrolyzing the linkage between a trehalose moiety and the remaining glycosyl moiety in a non-reducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher;
 - (2) Molecular weight
 - About 57,000 to 68,000 daltons on sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE):
 - (3) Isoelectric point (pl)
 - About 3.3 to 4.6 on isoelectrophoresis using ampholyte;
 - (4) Optimum temperature
 - About 35-45°C when incubated at pH 7.0 for 30 min;
 - (5) Optimum pH
 - About 6.0-7.5 when incubated at 40°C for 30 min: (6) Thermal stability

 - Stable up to a temperature of about 30-45°C when incubated at pH 7.0 for 60 min; and (7) pH Stability

 - Stable at a pH of about 5.0-10.0 when incubated at 25°C for 16 hours
 - 4. An enzyme according to any one of claims 1 to 3, which has one or more partial amino acid sequences selected from the group consisting of:
 - (1) leucine-aspartic acid-tryptophan-alanine-glutamic acid-alanine-X₁-X₂-glycine-aspartic acid (where X₁ means serine or alanine, and X₂ means alanine or glutamic acid);
 - (2) aspartic acid-glutamic acid-arginine-alanine-valine-histidine-isoleucine-leucine-glutamic acid-X3 (where X3 means glutamic acid or aspartic acid), and
 - (3) X₄-glycine-glutamic acid-glycine-asparagine-threonine-tryptophan-glycine-aspartic acid-serine (where X₄

means histidine or glutamine).

5

20

25

30

40

- 5. An enzyme according to any one of the preceding claims, which is an enzyme derived from a microorganism.
- An enzyme according to claim 5, wherein said microorganism is a microorganism selected from the group consisting of those of the genera Rhizobium, Arthrobacter, Brevibacterium and Micrococcus.
- An enzyme according to claim 6, wherein said microorganism of the genus Rhizobium is Rhizobium sp. M-11
 (FERM BP-4130)
 - An enzyme according to claim 6, wherein said microorganism of the genus Arthrobacter is Arthrobacter sp. Q36 (FERM BP-4316).
- 15 9. A process for proparing a trehalose-releasing enzyme according to any one of the preceding claims, said process comprising culturing a microgranism capable of producing said enzyme in a nutrient culture medium to form said enzyme, and recovering the resultant enzyme.
 - A process for preparing trehalose, comprising:
 - (a) allowing an enzyme to act on a solution containing a non-reducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher to form trehalose, said enzyme being capable of specifically hydrolyzing the linkage between a trehalose moiety and the remaining glycosyl moiety in said non-reducing saccharide;
 - (b) purifying the resultant trehalose; and
 - (c) recovering the purified trehalose.
 - 11. A process according to claim 10, wherein said enzyme in the step (a) is used together with a non-reducing saccharide-forming enzyme capable of forming one or more non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher.
 - 12. A process according to claim 10 or claim 11, wherein the step (a) further contains a step of allowing glucoamylase to act on the resultant solution in the step (a).
- 35 13. A process according to any one of claims 10 to 12, wherein the step (b) contains a step of subjecting the resultant solution containing trehalose in the step (a) to column chromatography using a column packed with a strong-acid cation-exchange resin to purily the trehalose.
 - 14. A process according to any one of claims 10 to 13, wherein the step (b) further contains a step of crystallizing trehalose in the resultant solution in the step (b) into hydrous- or anhydrous-crystalline trehalose.
 - 15. A process according to any one of claims 10 to 14, wherein said glycosyl moiety consists of one or more glucose residues
- 45 16. A process according to any one of claims 10 to 15, wherein said enzyme is as defined in any one of claims 3 to 8.
 - 17. A process for preparing saccharide composition containing trehalose, comprising:
 - (a) allowing an enzyme to act on a solution containing a non-reducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher to form trehalose, said enzyme being capable of specifically hydrolyzing the linkage between a trehalose moiety and the remaining glycosyl moiety in said non-reducing saccharide; and
 - (b) recovering the resultant saccharide composition containing trehalose and other saccharide(s).
- 18. A process according to claim 17. wherein said enzyme in the step (a) is used together with a non-reducing saccharide-forming enzyme capable of forming one or more non-reducing saccharides having a trehalose structure as an end unit and having a degree of ducese polymerization of 3 or higher.

- 19. A process according to claim 17 or claim 18, wherein the step (a) further contains a step of allowing glucoamylase to act on the resultant solution in the step (a).
- 20. A process according to any one of claims 17 to 19, wherein the step (a) further contains a step of crystallizing trehalose in the resultant solution in the step (a) into hydrous- or anhydrous-crystalline trehalose.
 - 21. A process according to any one of claims 17 to 20, wherein said glycosyl moiety consists of one or more glucose residues
- 10 22. A process according to any one of claims 17 to 21, wherein said enzyme in the step (a) is as defined in any one of claims 3 to 8.
 - 23. A process for preparing composition containing trehalose, comprising:
- 15 (a) allowing a non-reducing saccharide-forming enzyme (I) together with a trehalose-releasing enzyme (II) to act on a solution containing one or more reducing partial starch hydrolysates having a degree of glucose polymerization of 3 or higher to form trehalose, said enzyme (I) being capable of forming a non-reducing saccharide having a trehalose structure as an end unit and having a degree of glucose polymerization of 3 or higher, and said enzyme (II) being capable of specifically hydrolyzing the linkage between a trehalose moiety 20
 - and the remaining glycosyl moiety in said non-reducing saccharide;
 - (b) recovering the resultant trehalose together with or without other saccharide(s); and
 - (c) incorporating the trehalose together with or without other saccharide(s) into a material for composition.
- 24. A process according to claim 23, which produces a food product. 25
 - 25. A process according to claim 23, which produces a cosmetic composition.
 - 26. A process according to claim 23, which produces a pharmaceutical composition.
- 27. A process according to any one of claims 23 to 26, wherein said glycosyl moiety consists of one or more glucose residues.
 - 28. A process according to any one of claims 23 to 27, wherein said enzyme (II) is as defined in any one of claims 3 to 8.
- 35 29. A method to lower the degree of glucose polymerization of a reducing partial starch hydrolysate without increasing its reducing power, which contains a step of allowing a non-reducing saccharide-forming enzyme (I) together with a trehalose-releasing enzyme (II) to act on a solution containing one or more reducing partial starch hydrolysates having a degree of glucose polymerization of 3 or higher, said enzyme (I) being capable of forming one or more non-reducing saccharides having a trehalose structure as an end unit and having a degree of glucose polymeri-40 zation of 3 or higher, and said enzyme (II) being capable of specifically hydrolyzing the linkage between a trehalose molety and the remaining glycosyl molety in said non-reducing saccharide.
 - 30. A method according to claim 29, wherein said glycosyl moiety consists of one or more glucose residues.
- 31. A method according to claim 29 or claim 30, wherein said enzyme (II) is as defined in any one of claims 3 to 8.

Patentansprüche

- 50 1. Trehalose-freisetzendes Enzym, welches spezifisch die Bindung zwischen einer Trehalose-Komponente und der verbleibenden Glycosyl-Komponente in einem nicht-reduzierenden Saccharid mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder höher hydrolysiert.
- Enzym gemäß Anspruch 1, worin die Glycosyl-Komponente aus einem oder mehreren Glycoseresten besteht. 55
 - 3. Enzym gemäß Anspruch 1 oder Anspruch 2, welches die folgenden physikochemischen Eigenschaften aufweist:

(1) Wirkung

Spezifisches Hydrolysieren der Bindung zwischen einer Trehalose-Komponente und der verbleibenden Glycosyl-Komponente in einem nicht-reduzierenden Saccharid mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsprad von 3 oder höher:

(2) Molekulargewicht

Etwa 57000 bis 68000 Dalton mit Natriumdodecylsulfat-Polyacrylamid-Gelelektrophorese (SDS-PAGE); (3) Isoelektrischer Punkt (pl)

Etwa 3,3 bis 4,6 mit Isoelektrophorese unter Verwendung eines Ampholyten;

(4) Optimale Temperatur

Etwa 35-45°C, wenn man für 30 min bei pH 7,0 inkubiert;

(5) Optimaler pH

10

15

20

35

45

50

55

Etwa 6,0-7,5, wenn man für 30 min bei 40° inkubiert;

(6) Thermische Stabilität

Stabil bis zu einer Temperatur von etwa 30-45°C, wenn man für 60 min bei pH 7,0 inkubiert; und

(7) pH-Stabilität

Stabil bei einem pH von etwa 5,0-10,0, wenn man für 16 Stunden bei 25°C inkubiert.

- Enzym gemäß einem der Ansprüche 1 bis 3, welches eine oder mehrere Teil-Aminosäuresequenzen aufweist, gewählt aus der Gruppe, bestehend aus:
 - (1) Leucin-Asparaginsäure-Tryptophan-Alanin-Glutaminsäure-Alanin-X₁-X₂-Glycin-Asparaginsäure (worin X₁ Serin oder Alanin bedeutet und X₂ Alanin oder Glutaminsäure bedeutet);
 - (2) Asparaginsäure-Glutaminsäure-Arginin-Alanin-Valin-Histidin-Isoleucin-Leucin-Glutaminsäure-X₃ (worin
 - X₃ Glutaminsäure oder Asparaginsäure bedeutet); und
- (3) X₄-Glycin-Glutaminsäure-Glycin-Asparagin-Threonin-Tryptophan-Glycin-Asparaginsäure-Serin (worin X₄ Histidin oder Glutamin bedeutet).
 - Enzym gemäß einem der vorangehenden Ansprüche, welches ein aus einem Mikroorganismus gewonnenes Enzym ist.
- Enzym gem

 ß Anspruch 5, worin der Mikroorganismus ein Mikroorganismus ist, welcher aus der Gruppe, bestehend aus jenen der Gattungen Rhizobium, Arthrobacter, Brevibacterium und Micrococcus, gew

 ählt ist.
 - Enzym gemäß Anspruch 6, worin der Mikroorganismus der Gattung Rhizobium Rhizobium sp. M-11 (FERM BP-4130) ist.
 - Enzym gemäß Anspruch 6, worin der Mikroorganismus der Gattung Arthrobacter Arthrobacter sp. Q36 (FERM BP-4316) ist.
- Vorfahren zum Herstellen eines Trehalose-freisetzenden Enzyms gemäß einem der vorangehenden Ansprüche,
 wobei das Verfahren das Züchten eines Mikroorganismus, der das Enzym in einem N\u00e4hrkulturmed um zur Bildung des Enzyms produzieren kann, und das Gewinnen des resultierenden Enzyms umfaßt.
 - Verfahren zum Herstellen von Trehalose, umfassend:
 - (a) Einwirkenlassen eines Enzyms auf eine Lösung, die ein nicht-reduzierendes Saccharid mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder höher enthält, um Trehalose zu bilden, wobei das Enzym spezifisch die Bindung zwischen einer Trehalose-Komponente und der verbleibenden Glycosyl-Komponente in dem nicht-reduzierenden Saccharid hydrolysiseren kann;
 - (b) Reinigen der resultierenden Trehalose: und
 - (c) Gewinnen der gereinigten Trehalose.
 - 11. Verfahren gemäß Anspruch 10, worin das Enzym im Schritt (a) zusammen mit einem nicht-reduzierendes Saccharid bildenden Enzym verwendet wird, das ein oder mehrere nicht-reduzierende Saccharide mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder h\u00f6her bilden kann.
 - Verfahren gem

 ß Anspruch 10 oder Anspruch 11, worin der Schritt (a) weiters einen Schritt des Einwirkenlassens von Glucoamylase auf die resultierende Lösung im Schritt (a) umfaßt.

- 13. Verfahren gemäß einem der Ansprüche 10 bis 12, worin der Schritt (b) einen Schritt des Unterziehens der resultierenden Trehalose-enthaltenden Lösung im Schritt (a) einer Säulen-Chromatographie unter Verwendung einer Säule, die mit einem stark sauren Kationenaustauscherharz gepackt ist, umfaßt, um die Trehalose zu reinigen.
- 5 14. Verlahren gemäß einem der Ansprüche 10 bis 13, worin der Schritt (b) weiters einen Schritt des Kristallisierens von Trehalose in der resultierenden Lösung im Schritt (b) zu wasserhaltiger oder wasserfreier kristalliner Trehalose umfaßt
- 15. Verfahren gemäß einem der Ansprüche 10 bis 14, worin die Glycosyl-Komponente aus einem oder mehreren
 - Verfahren gemäß einem der Ansprüche 10 bis 15. worin das Enzym wie in einem der Ansprüche 3 bis 8 definiert ist.
 - 17. Verfahren zum Herstellen einer Trehalose enthaltenden Saccharid-Zusammensetzung, umfassend

15

20

25

- (a) Einwirkenlassen eines Enzyms auf eine Lösung, die ein nicht-reduzierendes Saccharid mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder h\u00f6her enth\u00e4lt, um Trehalose zu bilden, wobei das Enzym spezifisch die Bindung zwischen einer Trehalose-Komponente und der verbleibenden Glycosyl-Komponente in dem nicht-reduzierenden Saccharid hyddrolysieren kann: und
- (b) Gewinnen der resultierenden Saccharid-Zusammensetzung, die Trehalose und (ein) andere(s) Saccharid (e) enthält.
- 18. Verfahren gemäß Anspruch 17, worin das Enzym im Schritt (a) zusammen mit einem nicht-reduzierendes Saccharid bildenden Enzym verwendet wird, das ein oder mehrere nicht-reduzierende Saccharide mit einer Trehalose-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder h\u00f6her bilden kann.
- 20. Verfahren gemäß einem der Ansprüche 17 bis 19, worin der Schritt (a) welters einen Schritt des Kristallisierens von Trehalose in der resultierenden Lösung im Schritt (a) zu wasserhaltiger oder wasserfreiler kristalliner Trehalose umfaßt.
 - Verfahren gem
 äß einem der Anspr
 üche 17 bis 20, worin die Glycosyl-Komponente aus einem oder mehreren Glycoseresten besteht.
- 40 23. Verfahren zum Herstellen einer Trehalose enthaltenden Zusammensetzung, umfassend:
 - (a) Einwirkenlassen eines nicht-reduzierendes Saccharid bildenden Enzyms (I) zusammen mit einem Trehaleset-freisetzenden Enzym (II) auf eine Lösung, die ein oder mehrere reduzierende Stärke-Teilhydrolysate mit einem Glucose-Polymerisationsgrad von 3 oder höher enthält, im Trehalbes zu bilden, wobei das Enzym (I) ein nicht-reduzierendes Saccharid mit einer Trehalbes-Struktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder höher bilden kann, und das Enzym (II) spezifisch die Bilndung zwischen einer Trehalbes-Komponente in dem nicht-reduzierenden Saccharid hydrolysieren kann;
 - (b) Gewinnen der resultierenden Trehalose zusammen mit oder ohne (ein) andere(s) Saccharid(e): und
- (c) Einbauen der Trehalose zusammen mit oder ohne (ein) andere(s) Saccharid(e) in ein Zusammensetzungsmaterial.
 - 24. Verfahren gemäß Anspruch 23, worin ein Nahrungmittelprodukt hergestellt wird.
- Verfahren gemäß Anspruch 23, worin eine kosmetische Zusammensetzung hergestellt wird.
 - 26. Verfahren gemäß Anspruch 23, worin eine pharmazeutische Zusammensetzung hergestellt wird.

- Verfahren gemäß einem der Ansprüche 23 bis 26, worin die Glycosyl-Komponente aus einem oder mehreren Glucoseresten besteht
- 28. Verfahren gemäß einem der Ansprüche 23 bis 27, worin das Enzym (II) wie in einem der Ansprüche 3 bis 8 definiert
- 29. Verfahren zum Vermindern des Glucose-Polymerisationsgrades eines reduzierenden Stärke-Teilhydrolysals ohne Erhöhen seines Reduktionsvermögens, welches einen Schrift des Einwirkenlassens eines nicht-reduzierendes Saccharid blidenden Enzyms (I) zusammen mit einem Trehalose-freisetzenden Enzym (II) auf eine Lösung umfaßt, die ein oder mehrere reduzierendes Stärke-Teilhydrolysalte mit einem Glucose-Polymerisationsgrad von 3 oder höher erhalit, wobei das Enzym (I) ein oder mehrere nicht-reduzierende Saccharide mit einem Terhalose-Stuktur als Endeinheit und mit einem Glucose-Polymerisationsgrad von 3 oder höher bilden kann und das Enzym (II) spezilisch die Bindung zwischen einer Trehalose-Komponente und der verbleibenden Glycosyl-Komponente in dem nicht-reduzierenden Saccharid hydrolysieren kann.
- 30. Verfahren gemäß Anspruch 29. worin die Glycosyl-Komponente aus einem oder mehreren Glucoseresten besteht.
- Verfahren gemäß Anspruch 29 oder Anspruch 30, worin das Enzym (II) wie in einem der Ansprüche 3 bis 8 definiert ist

Revendications

10

15

25

30

35

40

45

50

- Enzyme libérant du tréhalose qui hydrolyse spécifiquement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans un saccharide non réducteur ayant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du glucose de 3 ou plus.
- Enzyme selon la revendication 1, dans laquelle ladite fraction glycosylique se compose d'un ou plusieurs résidus de glucose.
- 3. Enzyme selon la revendication 1 ou la revendication 2, qui a les propriétés physicochimiques suivantes :
 - (1) Action
 - Hydrolysant spécifiquement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans un saccharide non réducteur ayant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du quicose de 3 ou plus :
 - (2) Poids moléculaire
 - Environ 57 000 à 68 000 daltons par électrophorèse sur gel de polyacrylamide-dodécylsulfate de sodium (SDS-PAGE)
 - (3) Point isoélectrique (pl)
 - Environ 3,3 à 4,6 par isoélectrophorèse utilisant un ampholyte ;
 - (4) Température optimale
 - Environ 35 à 45°C en cas d'incubation à pH 7.0 pendant 30 min ;
 - (5) pH optimal
 - Environ 6,0 à 7,5 en cas d'incubation à 40°C pendant 30 min ;
 - (6) Stabilité thermique
 - Stable jusqu'à une température d'environ 30 à 45°C en cas d'incubation à pH 7,0 pendant 60 min | et (7) Stabilité au pH
 - Stable à un pH d'environ 5.0 à 10.0 en cas d'incubation à 25°C pendant 16 heures.
- 4. Enzyme selon l'une quelconque des revendications 1 à 3, qui a une ou plusieurs séquences partielles d'acides aminés choisies dans le groupe constitué par :
- (1) leucine-acide aspartique-tryptophane-alanine-acide glutamique-alanine-X₁-X₂-glycine-acide aspartique (où X₁ représente la sérine ou l'alanine, et X₂ représente l'alanine ou l'acide glutamique);
 - (2) acide aspartique-acide glutamique-arginine-alanine-valine-histidine-isoleucine-leucine-acide glutamique-X₂ (où X₂ représente l'acide glutamique ou l'acide aspartique); et
 - (3) X₄-glycine-acide glutamique-glycine-asparagine-thréonine-tryptophane-glycine-acide aspartique-sérine

- (où X₄ représente l'histidine ou la glutamine).
- Enzyme selon l'une quelconque des revendications précédentes, qui est une enzyme provenant d'un micro-organisme.
- Enzyme selon la revendication 5, dans laquelle ledit micro-organisme est un micro-organisme choisi dans le groupe constitué par ceux des genres Rhizobium, Arthrobacter, Brevibacterium et Micrococcus.
- Enzyme selon la revendication 6, dans laquelle ledit micro-organisme du genre Rhizobium est le Rhizobium sp.
 M-11 (FERM BP-4130).
 - Enzyme selon la revendication 6, dans laquelle ledit micro-organisme du genre Arthrobacter est l'Arthrobacter sp. Q36 (FERM BP-4316).
- 15 9. Procédé de préparation d'une enzyme libérant du tréhalces seton l'une quelconque des revendications précédentes, ledit procédé comportant la culture d'un micro-organisme capable de produire ladite enzyme dans un milieu de culture nutritif pour former ladite enzyme, et la récupération de l'enzyme résultante
 - 10. Procédé de préparation du tréhalose, comportant :
 - (a) l'action d'une enzyme sur une solution contenant un saccharide non réducteur ayant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du glucose de 3 ou plus pour former du tréhalose, ladite enzyme étant capable d'hydrolyser spécifiquement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans ledit saccharide non réducteur;
- 25 (b) la purification du tréhalose résultant ; et

20

30

40

45

50

- (c) la récupération du tréhalose purifié.
- 11. Procédé selon la revendication 10, dans lequel ladite enzyme de l'étape (a) est utilisée avec une enzyme formant un saccharide non réducteur capable de former un ou pluisieurs saccharides non réducteurs ayant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du glucose de 3 ou plus.
 - 12. Procédé selon la revendication 10 ou la revendication 11, dans lequel l'étape (a) contient en outre une étape consistant à faire agir de la glucoamylase sur la solution résultante dans l'étape (a).
- 35 13. Procédé selon l'une quelconque des revendications 10 à 12. dans lequel l'étape (b) contient une étape de soumission de la solution résultante contenant du tréhelose de l'étape (a) à la chromatographie sur colonne utilisant une colonne garnie d'une résine échangeuse de cations fortement acides pour purifier le tréhalose.
 - 14. Procédé selon l'une quelconque des revendications 10 à 13, dans lequel l'étape (b) contient en outre une étape de cristallisation du tréhalose de la solution résultante dans l'étape (b) en tréhalose cristallin hybrieté ou en tréhalose cristallin anhydre.
 - 15. Procédé selon l'une quelconque des revendications 10 à 14, dans lequel ladite fraction glycosylique se compose d'un ou plusieurs résidus de glucose.
 - 16. Procédé selon l'une quelconque des revendications 10 à 15, dans lequel ladite enzyme est définie comme indiqué dans l'une quelconque des revendications 3 à B.
 - 17. Procédé de préparation d'une composition de saccharides contenant du tréhalose, comportant :
 - (a) l'action d'une enzyme sur une solution contenant un saccharide non réducteur ayant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du glucose de 3 ou plus pour former du tréhalose, ladite enzyme étant capable d'hydrolyser spécifiquement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans ledit saccharide non réducteur; et
 - (b) la récupération de la composition de saccharides résultante contenant du tréhalose et autre(s) saccharide
 - 18. Procédé selon la revendication 17, dans lequel ladite enzyme dans l'étape (a) est utilisée avec une enzyme formant

un saccharide non réducteur capable de former un ou plusieurs saccharides non réducteurs avant une structure tréhalosique comme unité terminale et ayant un degré de polymérisation du glucose de 3 ou plus

- 19. Procédé selon la revendication 17 ou la revendication 18, dans lequel l'étape (a) contient en outre une étape consistant à faire agir de la glucoamylase sur la solution résultante dans l'étape (a).
- 20. Procédé selon l'une quelconque des revendications 17 à 19, dans lequel l'étape (a) contient en outre une étape de cristallisation du tréhalose de la solution résultante dans l'étape (a) en tréhalose cristallin hydraté ou en tréhalose cristallin anhydre.
- 21. Procédé selon l'une quelconque des revendications 17 à 20, dans lequel ladite fraction glycosylique se compose d'un ou plusieurs résidus de alucose.
- 22. Procédé selon l'une quelconque des revendications 17 à 21, dans lequel ladite enzyme de l'étape (a) est définie comme indiqué dans l'une quelconque des revendications 3 à 8.
- 23. Procédé de préparation d'une composition contenant du tréhalose, comportant :
- (a) l'action d'une enzyme formant un saccharide non réducteur (I) avec une enzyme libérant du tréhalose (II) 20 sur une solution contenant un ou plusieurs hydrolysats partiels d'amidon réducteur avant un degré de polymérisation du glucose de 3 ou plus pour former du tréhalose, ladite enzyme (I) étant capable de former un saccharide non réducteur avant une structure tréhalosique comme unité terminale et avant un degré de polymérisation du glucose de 3 ou plus, et ladite enzyme (II) étant capable d'hydrolyser spécifiquement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans ledit saccharide non réducteur ; 25
 - (b) la récupération du tréhalose résultant avec ou sans autre(s) saccharide(s) ; et
 - (c) l'incorporation du tréhalose avec ou sans autre(s) saccharide(s) dans une matière pour composition.
 - 24. Procédé selon la revendication 23, qui fournit un produit alimentaire.
- 25. Procédé selon la revendication 23, qui fournit une composition cosmétique.
 - 26. Procédé selon la revendication 23, qui fournit une composition pharmaceutique
- 27. Procédé selon l'une quelconque des revendications 23 à 26, dans lequel ladite fraction glycosylique se compose 35 d'un ou plusieurs résidus de alucose.
 - 28. Procédé selon l'une quelconque des revendications 23 à 27, dans lequel ladite enzyme (II) est définie comme indiqué dans l'une quelconque des revendications 3 à 8.
- 29. Procédé pour réduire le degré de polymérisation du glucose d'un hydrolysat partiel d'amidon réducteur sans augmenter son pouvoir réducteur, qui contient une étape d'action d'une enzyme formant un saccharide non réducteur (I) avec une enzyme libérant du tréhalose (II) sur une solution contenant un ou plusieurs hydrolysats partiels d'arnidon réducteur ayant un degré de polymérisation du glucose de 3 ou plus, ladite enzyme (I) étant capable de former un ou plusieurs saccharides non réducteurs ayant une structure tréhalosique comme unité terminale et 45 avant un degré de polymérisation du glucose de 3 ou plus, et ladite enzyme (II) étant capable d'hydrolyser spécifiguement la liaison entre une fraction tréhalosique et la fraction glycosylique restante dans ledit saccharide non réducteur
- 30. Procédé selon la revendication 29, dans lequel ladite fraction alvoosylique se compose d'un ou plusieurs résidus 50 de alucose.
 - 31. Procédé selon la revendication 29 ou la revendication 30, dans lequel ladite enzyme (II) est définie comme indiqué dans l'une quelconque des revendications 3 à 8.

10

















